

# ASCEND IV

## Advanced System for Computations in ENgineering Design

A portable mathematical modeling environment.

Release 0.8

Sept 26, 1997

### Authors:

The research group of Arthur Westerberg  
Department of Chemical Engineering, Carnegie Mellon University  
Benjamin Allan, Vicente Rico-Ramirez, Mark Thomas, Kenneth Tyner  
Other Helpful People Too Numerous To Mention Here

### Sponsors 1993-1997:

Computer Aided Process Design Consortium  
Institute for Complex Engineered Systems  
National Science Foundation grant for the Engineering Design Research Center  
US Department of Energy

We are deeply indebted to the authors and contributors at large who created Tcl/Tk. Many thanks  
Dr. Osterhout!

### Keywords:

ASCEND, CONOPT, EDRC, FORTRAN, GAMS, GNU license, GUI, ICES, LSODE, Levenberg-Marquardt, MINLP, NLP, Newton, ODE, Tcl, Tk, UNIFAC, boundary value, chemical engineering, collocation, complex engineered system,, conditional modeling, copyleft, degrees of freedom, design, design research center, distillation, dynamic, engineering design, free software, freeware, initial value, initialization, interactive, large-scale, linear algebra, linear equations, mathematical modeling, mixed integer, modeling system, nonideal thermodynamics, nonlinear program, object oriented, optimization, ordinary differential equation, Pitzer vapor, reactive distillation, scalable, scaling, simulation, solving, structural analysis, Wilson liquid.

## Documentation Bird's Eye View

ASCEND IV	1
Advanced System for Computations in ENgineering Design	1
Documentation Bird's Eye View	2
Documentation Detail Map	4
A typical scenario for running the ASCEND system	16
Getting Started with ASCEND	20
Script	22
Library	32
Merged into library	42
Browser	46
Solver	56
The Data Probe Window	68
ASC PLOT	74
Display slave	82
ASCEND Units	84
The ASCEND Toolbox	88
The System Utilities Window	92
Font Selection Dialog	100
The Print Dialog	104
Solved simple modeling problems with ASCEND	108
A Conditional Modeling Example: Representing a Superstructure	114
A Simple Chemical Engineering Flowsheeting Example	138
The ASCEND predefined collection of models	156
The ASCEND IV language syntax and semantics	158
Units library	218
Brief History of ASCEND	232



## Documentation Detail Map

<b>A typical scenario for running the ASCEND system</b>	<b>16</b>
<b>Getting Started with ASCEND</b>	<b>20</b>
:: Philosophy	20
- Getting the ASCEND system and installing it	20
- Starting ASCEND	21
o ASCENDDIST	21
o ASCENDHELP	21
o ASCENDLIBRARY	21
<b>Script</b>	<b>22</b>
Figure ASCEND's Script Window.	22
:: The Script Menu Bar	23
- Script File Menu	23
o New File	23
o Read File	23
o Import File	23
o Save	23
o Save As	23
o Buffer List	23
- Script Edit Menu	23
o Write selection	23
o Select all	23
o Remove statements	23
- Script Execute Menu	24
o Statements selected	24
- Script toolbox menu	24
- Script Help menu	24
o On ASCEND/TCL Scripts	24
o On SCRIPT	24
:: SCRIPT Grill Menu	24
o Record actions	24
:: The Script Language	24
- Summary	24
o <arg>	25
o <a1,a2>	25
o <a1 a2>	25
o [a1]	25
o [a,b]	25
o qlfdid	25
o qlfpid	25
o {}	25
- Quick reference:	25
o ASSIGN	25
o BROWSE	25

	<i>o</i>	<i>CLEAR_VARS</i>	25
	<i>o</i>	<i>COMPILE</i>	25
	<i>o</i>	<i>DELETE</i>	25
	<i>o</i>	<i>DISPLAY*</i>	25
	<i>o</i>	<i>INTEGRATE</i>	25
	<i>o</i>	<i>MERGE</i>	25
	<i>o</i>	<i>PLOT</i>	26
	<i>o</i>	<i>PRINT</i>	26
	<i>o</i>	<i>PROBE</i>	26
	<i>o</i>	<i>READ</i>	26
	<i>o</i>	<i>REFINE</i>	26
	<i>o</i>	<i>RESTORE*</i>	26
	<i>o</i>	<i>RESUME</i>	26
	<i>o</i>	<i>RUN</i>	26
	<i>o</i>	<i>SAVE*</i>	26
	<i>o</i>	<i>SHOW</i>	26
	<i>o</i>	<i>SOLVE</i>	26
	<i>o</i>	<i>WRITE</i>	26
-		Commands	26
	<i>o</i>	<i>ASSIGN</i>	26
	<i>o</i>	<i>BROWSE</i>	26
	<i>o</i>	<i>CLEAR_VARS</i>	26
	<i>o</i>	<i>COMPILE</i>	26
	<i>o</i>	<i>DELETE</i>	27
	<i>o</i>	<i>DISPLAY</i>	27
	<i>o</i>	<i>INTEGRATE</i>	27
	<i>o</i>	<i>MERGE</i>	27
	<i>o</i>	<i>OBJECTIVE</i>	27
	<i>o</i>	<i>PLOT</i>	27
	<i>o</i>	<i>PRINT</i>	28
	<i>o</i>	<i>PROBE</i>	28
	<i>o</i>	<i>READ</i>	28
	<i>o</i>	<i>REFINE</i>	28
	<i>o</i>	<i>RESTORE</i>	28
	<i>o</i>	<i>RESUME</i>	29
	<i>o</i>	<i>RUN</i>	29
	<i>o</i>	<i>SAVE</i>	29
	<i>o</i>	<i>SHOW</i>	29
	<i>o</i>	<i>SOLVE</i>	29
	<i>o</i>	<i>WRITE</i>	29
	<i>o</i>		29
::		Script Window Bindings	29
	<i>o</i>	<i>M1</i>	30
	<i>o</i>	<i>M1-Drag</i>	30
	<i>o</i>	<i>Shift-M1[-Drag]</i>	30
	<i>o</i>	<i>Double-M1</i>	30

	<i>o Double-M1-Drag</i>	30
	<i>o Triple-M1</i>	30
	<i>o Triple-M1-Drag</i>	30
	<i>o M2</i>	30
	<i>o M2-Held-Down</i>	30
	<i>o M3</i>	30
	<i>o Control-M1</i>	30
	<i>o Control-k</i>	30
	<i>o Control-w</i>	30
	<i>o Meta-w</i>	30
	<i>o Control-y</i>	30
	<i>o Meta-y</i>	30
<b>Library</b>		<b>32</b>
	Figure ASCEND Library Window.	32
	Figure Data structure used to store type defini-	
tions.		33
::	Menu Bar	34
-	The file Menu	34
	<i>o Read types from file</i>	34
	<i>o Close window</i>	34
-	The Edit Menu	34
	<i>o Create simulation</i>	34
	<i>o Delete Simulation</i>	35
	<i>o Delete all types</i>	35
	Figure The Create Simulation Dialog	35
-	The Display Menu	35
	<i>o Code</i>	35
	<i>o Ancestry</i>	35
	<i>o Refinement hierarchy</i>	35
	<i>o External Functions</i>	35
	<i>o Hide Type</i>	35
	<i>o UnHide Type</i>	35
	<i>o Hide/Show Fundamentals</i>	36
	Figure Select the fundamental type to Hide or	
Unhide.		36
-	The Find Menu	36
	<i>o Type by name</i>	36
	Figure The Library's Find Type dialog.	36
	<i>o Type by fuzzy name</i>	37
	<i>o Pending statements</i>	37
	<i>o To Display</i>	37
	<i>o To Console</i>	37
	<i>o To File</i>	37
-	The View Menu	37
-	The export Menu	37
	<i>o Simulation to Browser</i>	37

	o	<i>Simulation to Solver</i>	37
	o	<i>Simulation to Probe</i>	38
-		The help Menu	38
	o	<i>On LIBRARY</i>	38
::		Type Refinement Hierarchy Window	38
		Figure The Type Refinement Window.	38
		Figure The Parts window displays the parts.	39
		Figure The Hierarchy Roots Window.	40
		<b>Merged into library</b>	<b>42</b>
::		Sims Window	42
		Figure	42
		Figure	43
-		The Edit menu	43
-		The Pendings menu	44
-		The Export menu	44
		<b>Browser</b>	<b>46</b>
		Figure ASCEND's Browser window.	46
::		The Menu Bar	47
-		BROWSER Edit Menu	47
	o	<i>Run method</i>	47
	o	<i>Clear Vars</i>	47
	o	<i>Set value</i>	47
	o	<i>Read values</i>	48
	o	<i>Refine</i>	48
	o	<i>Merge</i>	48
	o	<i>Compile</i>	48
	o	<i>Resume Compilation</i>	48
	o	<i>Create Part</i>	48
-		BROWSER Display menu	49
	o	<i>Attributes</i>	49
	o	<i>Relations</i>	49
	o	<i>Cond Rels</i>	49
	o	<i>Log Rels</i>	49
	o	<i>Cond Log Rels</i>	49
	o	<i>Whens</i>	49
	o	<i>Plot</i>	50
	o	<i>Statistics</i>	50
-		BROWSER Find menu	50
	o	<i>By name</i>	50
	o	<i>By type</i>	50
	o	<i>Aliases</i>	52
	o	<i>Where created</i>	52
	o	<i>Clique</i>	52
	o	<i>Eligible variables</i>	52
	o	<i>Relations</i>	53
	o	<i>Operands</i>	53

	o	Parents	53
	o	Pendings	53
-		BROWSER view menu	53
	o	Suppress Atoms	53
	o	Display Atom Values	53
	o	Check Dimensionality	53
	o	Hide Names	53
	o	UnHide Names	53
-		BROWSER Export menu	53
	o	to Solver	53
	o	Many to Probe	54
		Figure Filtering instances sent to the Probe	54
	o	Item to Probe	54
-		BROWSER Help menu	54
	o	On BROWSER	54
		<b>Solver</b>	<b>56</b>
		Figure Solver Window	56
::		The Solver Menu Bar	57
-		Solver Edit Menu	57
	o	Remove instance	57
	o	Select objective	57
-		Solver Display Menu	57
	o	Status	57
	o	Unattached variables	57
	o	Unincluded relations	57
	o	Incidence matrix	57
		Figure The Incidence Matrix	58
-		Solver Execute Menu	58
	o	Solve	58
	o	Single step	58
	o	Integrate	58
-		Solver Analyze menu	58
	o	Reanalyze	58
	o	Debugger	58
	o	Overspecified	59
	o	Find dependent eqns.	59
	o	Find unassigned eqns.	59
	o	Evaluate unincluded eqns.	59
	o	Find vars near bounds	59
	o	Find vars far from nom	59
-		Solver Export Menu	60
	o	to Browser	60
	o	to Probe	60
::		Solver Button Bar	60
	o	Solver Select Button	60
	o	Solver Options Button	60



o	Halt Button	60
-	General parameters page	60
	Figure General Parameter Page	61
::	Available Solvers	62
-	QRSlv	63
::	Debugger	64
	Figure The Debugger Window	65
	<b>The Data Probe Window</b>	<b>68</b>
::	Overview	68
	Figure Probe window	68
::	The File menu	69
-	New buffer	69
-	Read file	69
-	Save	69
-	Save as	70
-	Print	70
::	The Edit Menu	70
-	Remove Selected names	70
-	Remove all names	70
-	Remove UNCERTAIN names	70
-	Copy	70
::	The View Menu	70
::	The Export Menu	71
-	to Browser	71
-	to Display	71
::	The Probe Filter	71
	Figure Probe import filter	72
	<b>ASC PLOT</b>	<b>74</b>
::	Plot maker	74
	Figure The Ascend Plot Window	74
-	The Edit Menu	75
-	The Execute Menu	75
	Figure The Create Data Window	76
-	The Display Menu	77
	Figure The Graph Generics Window	78
	Figure Complete Plot	79
::	Navigation	80
	Figure Phase Diagram	81
	<b>Display slave</b>	<b>82</b>
::	Overview	82
	Figure Display slave window	82
::	The File Menu	83
-	Print	83
-	Close window	83
::	The View Menu	83
-	Show comments in code	83

-	Font	83
-	Open automatically	83
::	Title line	83
	<b>ASCEND Units</b>	<b>84</b>
::	The Menu Bar	84
	<i>o</i> Units vs dimensions	84
	<i>o</i> Typical use	84
-	UNITS Edit Menu	85
	<i>o</i> Set precision	85
	<i>o</i> Read file	85
	<i>o</i> Write file	85
-	UNITS Display Menu	85
	<i>o</i> Show all units	85
	<i>o</i> SI(MKS)	85
	<i>o</i> US Engineering	85
	<i>o</i> CGS	85
-	UNITS Help Menu	85
	<i>o</i> An essay on units vs dimensions	85
	<i>o</i> On UNITS	86
	<b>The ASCEND Toolbox</b>	<b>88</b>
	Figure The ASCEND Toolbox window.	88
::	Exit	89
::	Ascplot	89
::	Help	89
::	Utilities	89
::	Bug Report	89
	<b>The System Utilities Window</b>	<b>92</b>
::	Overview	92
	Figure The System Utilities window manages ASCEND's interaction with the operating system and with other programs.	92
::	Variables	93
-	WWW Root URL	93
-	WWW Restart Command	94
-	WWW Startup Command	94
-	ASCENDLIBRARY Path*	94
-	Scratch Directory	95
-	Working Directory	95
-	Plot Program Type	95
-	Plot Program Name	95
-	Text Edit Command	95
-	Postscript Viewer	96
-	Spreadsheet Command	96
-	Text Print Command	96
-	PRINTER Variable*	96
-	ASCENDDIST Directory*	96

-	TCL_LIBRARY Environment Variable*	97
-	TK_LIBRARY Environment Variable*	97
::	Buttons	97
-	OK	98
-	Save	98
-	Read	98
-	More	98
-	Help	98
	<b>Font Selection Dialog</b>	<b>100</b>
::	Overview	100
	Figure The font selection dialog.	100
::	Font Menu	101
::	Style Menu	101
::	Cancel Button	101
::	OK Button	101
::	Current Font Sample	102
::	Font Sampler Area	102
::	Point Size Slider	102
::	Current Font Selection	102
::	Setting the Default Font	102
	<b>The Print Dialog</b>	<b>104</b>
::	Overview	104
	Figure The print dialog.	104
::	Settings	104
-	Destination	104
-	Printer	106
-	Name of file	106
-	Enscript flags	106
-	User print command	106
::	Buttons	107
-	OK	107
-	Help	107
-	Cancel	107
	<b>Solved simple modeling problems with ASCEND</b>	<b>108</b>
::	Roots of a polynomial	108
-	Problem statement	109
-	Answer	109
::	Numerical integration of tabular data	110
-	Problem statement	110
-	Answer	111
	<b>A Conditional Modeling Example: Representing a Superstructure</b>	<b>114</b>
	Figure Superstructure used in the example of the application of the when statement	114
::	The WHEN Statement	114
::	The Problem Description	116

::	The Code	116
	<b>A Simple Chemical Engineering Flowsheeting Example</b>	<b>138</b>
::	The problem description	138
::	The code	139
	<b>The ASCEND predefined collection of models</b>	<b>156</b>
	<i>o</i> <i>system.lib</i>	156
	<i>o</i> <i>atoms.lib</i>	156
	<i>o</i> <i>Typical use of library files</i>	157
	<i>o</i> <i>Examples and scripts</i>	157
	<b>The ASCEND IV language syntax and semantics</b>	<b>158</b>
::	Preliminaries	159
-	Punctuation	160
	<i>o</i> <i>keywords:</i>	160
	<i>o</i> <i>(* *)</i>	161
	<i>o</i> <i>()</i>	161
	<i>o</i> <i>{ }</i>	161
	<i>o</i> <i>[ ]</i>	162
	<i>o</i> <i>.</i>	162
	<i>o</i> <i>..</i>	162
	<i>o</i> <i>:</i>	162
	<i>o</i> <i>::</i>	162
	<i>o</i> <i>;</i>	162
-	Basic Elements	162
	<i>o</i> <i>L</i>	163
	<i>o</i> <i>M</i>	163
	<i>o</i> <i>T</i>	163
	<i>o</i> <i>E</i>	163
	<i>o</i> <i>Q</i>	164
	<i>o</i> <i>TMP</i>	164
	<i>o</i> <i>LUM</i>	164
	<i>o</i> <i>P</i>	164
	<i>o</i> <i>S</i>	164
	<i>o</i> <i>C</i>	164
-	Basic Concepts	169
::	Data Type Declarations	172
	<i>o</i> <i>UNIVERSAL</i>	172
-	Models	173
	<i>o</i> <i>MODEL</i>	173
	<i>o</i> <i>foo</i>	173
	<i>o</i> <i>bar</i>	173
	<i>o</i> <i>column(n,s)</i>	173
	<i>o</i> <i>flowsheet</i>	174
-	Sets	174
	<i>o</i> <i>:=</i>	174
	<i>o</i> <i>UNION[setlist]</i>	175
	<i>o</i> <i>+</i>	175

	<i>o</i>	<i>INTERSECTION[]</i>	175
	<i>o</i>	<i>*</i>	175
	<i>o</i>	<i>-</i>	175
	<i>o</i>	<i>CARD[set]</i>	175
	<i>o</i>	<i>CHOICE[set]</i>	175
	<i>o</i>	<i>IN</i>	176
	<i>o</i>	<i>SUCH_THAT (* 4 *)</i>	176
	<i>o</i>	<i>/</i>	177
-		Constants	177
	<i>o</i>	<i>real_constant</i>	177
	<i>o</i>	<i>integer_constant</i>	178
	<i>o</i>	<i>symbol_constant</i>	178
	<i>o</i>	<i>boolean_constant</i>	178
	<i>o</i>	<i>:=</i>	178
-		Variables	178
	<i>o</i>	<i>ATOM</i>	178
	<i>o</i>	<i>DEFAULT, DIMENSION, and DIMEN-</i>	
<i>SIONLESS</i>		179	
	<i>o</i>	<i>real</i>	179
	<i>o</i>	<i>integer</i>	179
	<i>o</i>	<i>boolean</i>	179
	<i>o</i>	<i>symbol</i>	180
	<i>o</i>	<i>:=</i>	180
	<i>o</i>	<i>DATA (* 4+ *)</i>	180
	<i>o</i>		181
-		Relations	181
	<i>o</i>	<i>=, &gt;=, &lt;=, &lt;, &gt;, &lt;&gt;</i>	182
	<i>o</i>	<i>MAXIMIZE, MINIMIZE</i>	182
	<i>o</i>	<i>+</i>	182
	<i>o</i>	<i>-</i>	182
	<i>o</i>	<i>*</i>	182
	<i>o</i>	<i>/</i>	182
	<i>o</i>	<i>^</i>	183
	<i>o</i>	<i>-</i>	183
	<i>o</i>	<i>ordered_function()</i>	183
	<i>o</i>	<i>SUM[term set]</i>	183
	<i>o</i>	<i>PROD[term set]</i>	183
	<i>o</i>	<i>MAX[term set]</i>	184
	<i>o</i>	<i>MIN[term set]</i>	184
-		Derivatives in relations (* 4+ *)	184
-		External relations	184
-		Conditional relations (* 4 *)	184
-		Logical relations (* 4 *)	184
-		NOTES (* 4+ *)	185
::		Declarative statements	185
	<i>o</i>	<i>IS_A</i>	186

	<i>o</i>	<i>IS_REFINED_TO</i>	186
	<i>o</i>	<i>ALIASES (* 4 *)</i>	186
	<i>o</i>	<i>ALIASES/ISA (*4*)</i>	187
	<i>o</i>	<i>WILL_BE (* 4 *)</i>	187
	<i>o</i>	<i>ARE_THE_SAME</i>	187
	<i>o</i>	<i>WILL_BE_THE_SAME (* 4 *)</i>	187
	<i>o</i>	<i>WILL_NOT_BE_THE_SAME (* 4 *)</i>	187
	<i>o</i>	<i>ARE_NOT_THE_SAME (* 4+ *)</i>	187
	<i>o</i>	<i>ARE_ALIKE</i>	187
	<i>o</i>	<i>FOR/CREATE</i>	187
	<i>o</i>	<i>SELECT/CASE (*4*)</i>	188
	<i>o</i>	<i>CONDITIONAL (*4*)</i>	188
	<i>o</i>	<i>WHEN/CASE (* 4 *)</i>	188
	<i>o</i>	<i>IS_A</i>	188
	<i>o</i>	<i>IS_REFINED_TO</i>	189
	<i>o</i>	<i>ALIASES (* 4 *)</i>	189
	<i>o</i>	<i>ALIASES/ISA (*4*)</i>	190
	<i>o</i>	<i>WILL_BE (* 4 *)</i>	191
	<i>o</i>	<i>ARE_THE_SAME</i>	191
	<i>o</i>	<i>WILL_BE_THE_SAME (* 4 *)</i>	193
	<i>o</i>	<i>WILL_NOT_BE_THE_SAME (* 4 *)</i>	193
	<i>o</i>	<i>ARE_NOT_THE_SAME (* 4+ *)</i>	194
	<i>o</i>	<i>ARE_ALIKE</i>	194
	<i>o</i>	<i>FOR/CREATE</i>	195
	<i>o</i>	<i>SELECT/CASE (*4*)</i>	195
	<i>o</i>	<i>CONDITIONAL (*4*)</i>	196
	<i>o</i>	<i>WHEN/CASE (* 4 *)</i>	196
<i>::</i>		Procedural statements	196
	<i>o</i>	<i>METHODS</i>	196
	<i>o</i>	<i>ADD METHODS IN type_name; (*4+*)</i>	
196			
	<i>o</i>	<i>REPLACE METHODS IN type_name;</i>	
(*4+*)		197	
	<i>o</i>	<i>METHOD</i>	197
	<i>o</i>	<i>FOR/DO statement</i>	198
	<i>o</i>	<i>IF</i>	198
	<i>o</i>	<i>SWITCH (* 4+ *)</i>	199
	<i>o</i>	<i>CALL</i>	199
	<i>o</i>	<i>RUN</i>	199
<i>::</i>		Parameterized models	199
-		The parameter list	200
-		The WHERE list	201
-		The assignment list	201
-		Refining parameterized types	201
<i>::</i>		Miscellany	202
-		Variables for solvers	202

	<i>o solver_var</i>	202
	<i>o lower_bound</i>	203
	<i>o upper_bound</i>	203
	<i>o nominal</i>	203
	<i>o fixed</i>	203
	<i>o generic_real</i>	203
	<i>o solver_semi, solver_integer,</i>	
<i>solver_binary</i>	203	
	<i>o ivpsystem.lib</i>	204
-	Supported attributes	204
	<i>o (* 4+ *)</i>	204
-	Single operand real functions:	204
	<i>o exp()</i>	204
	<i>o ln()</i>	204
	<i>o sin()</i>	204
	<i>o cos()</i>	204
	<i>o tan()</i>	204
	<i>o arcsin()</i>	205
	<i>o arccos()</i>	205
	<i>o arctan()</i>	205
	<i>o erf()</i>	205
	<i>o sinh()</i>	205
	<i>o cosh()</i>	205
	<i>o tanh()</i>	205
	<i>o arcsinh()</i>	205
	<i>o arccosh()</i>	205
	<i>o arctanh()</i>	205
	<i>o lnm()</i>	205
	<i>o abs()</i>	206
-	Logical functions	206
	<i>o SATISFIED() (*4*)</i>	206
-	UNITS definitions	206
	<b>Units library</b>	<b>218</b>
::	Units	218
::	The basic units in an extended SI MKS system	218
::	Units defined in measures.a4l, the default system units library of atoms.a4l.	219

# CHAPTER 1 A TYPICAL SCENARIO FOR RUNNING THE ASCEND SYSTEM

The ASCEND system is a modeling environment. We have designed it to allow modelers to pose, debug and solve or optimize models described by up to of the order of a hundred thousand nonlinear algebraic equations on a conventional UNIX workstation or PC running Windows NT4 or Windows 95. The ASCEND system comprises three major parts: the ASCEND modeling language for posing models, the ASCEND interactive environment to allow users to compile, debug and execute models, and a suite of solvers and optimizers.

You would typically proceed as follows to use the ASCEND system for modeling.

1. Using your favorite text editor (e.g., xemacs<sup>1</sup>), you will create a model of the problem you wish to solve in the ASCEND modeling language. ASCEND models are type definitions. Each model typically includes a declaration of the parts from which it is constructed, including variables, instances of previously defined types and arrays of any of these. Each model also includes the equations it adds to the model definition over and above those equations that its parts will provide. Finally if you abide by our advice on model writing, you will also write three or four methods that you will later run interactively on the compiled model instance to prepare it to be solved.

If the model is particularly complex, you will probably create your model using types defined earlier by yourself and others. For chemical process flowsheet models, we provide a library of types. We also provide a file that contains most of the types of variables and constants anyone would use to construct a model.

2. Start up the interactive ASCEND user interface by typing 'ascend' on the Unix shell command line or double clicking on the ASCEND icon on a PC. A number of different tool sets, each represented by a special window, open up on the screen. The one you will likely focus on at first is the SCRIPT window. In this

---

1. Xemacs is a very powerful text editor which is widely used on UNIX workstations. It is available for free for both UNIX workstations and PCs through the WWW (search on xemacs).



window under Tools, you will open whichever tool window you want to work with at first, probably the LIBRARY tool set which provides tools to load text files containing ASCEND models. You will likely move and/or resize these windows.

3. Using a tool in the LIBRARY tool set, you will load the files containing the previously defined types of which your model makes use. You will open last the file containing the model you have just written. Unless you are incredibly skilled and/or lucky, you will see several error messages indicating that you have not correctly posed your model as the system attempts to load your new file. Moving back to your favorite editor you will correct syntax errors discovered by this file loading process, attempt again to load, make more corrections, etc.
4. Once your new model description can pass the loading process without errors, you will compile a simulation for it. Again there could be errors.
5. You will export this compiled simulation to the BROWSER tool set so you can look at the model, examining all its parts. If there were compiler errors, you may use tools in this tool set to aid you to find exactly what you have done wrong in posing the model.

For example, if it is a particularly complex model, you will methodically examine it to see if you have configured it as you wanted.

6. Once you are through inspecting the model and have removed all the errors that of which you are aware, you will prepare the model to be solved. This you will do by asking the system to execute the methods you should have written to go with your model description. If you abide by the style of modeling that we strongly advocate, your model will have these methods attached to it -- written before your first attempt to compile it. These methods will be for setting initial values for the variables, for scaling the variables, and for setting the "fixed" flags for a sufficient number of variables to make the model instance well-posed. To be well-posed means a number of things, among them that the model has the same number of variables to be calculated as equations available for calculating them.
7. You will next export the model to the SOLVER tool set. When importing a model, the SOLVER tool set analyzes the model to discover how many variables and equations are in its description. If it is not an optimization problem, the SOLVER looks to see if it is well-posed and, if not, will issue warning messages and open up an interactive tool provided to aid you to make it well-posed right then and there. What you learn while using this tool you

will likely encode right away into the model description so the next time you compile this model, it will become well-posed without this interactive step.

8. You can interactively choose among the available solvers and will most likely choose our nonlinear equation solving solver. With fingers crossed, you will ask the solver to start solving.
9. Whether or not it solves successfully, you will likely return to the BROWSER to inspect the results as you can view the value for every variable and equation residual in the model using the BROWSER. If the solving process fails, you can select tools both in the BROWSER and the SOLVER to look for the likely problem. For example, you may have posed your problem and its initial conditions such that the solution is out of bounds. A tool will tell you if the SOLVER has driven any of the variables in the model to their bounds. Another will tell you if some of your variables are poorly scaled. Yet another will investigate the model to see if it is locally singular, and if it is, that tool will report to you exactly which equations (by name) have given it reason to believe that to be so. (In the near future, this tool will also tell you that you should change what you are fixing and what you are calculating to remove this singularity, if such a move would prove useful. It will give you a list of variables from which to choose for each of these trades.)
10. You may wish to see the output in units different from those currently used. Opening the UNITS tools set will allow you to change wholesale from SI to American Engineering and/or to change individual units such as those for pressure from bars to atm.
11. You may have opened the SCRIPT tool set before loading the model files. Before doing all the above steps within ASCEND, you may then have activated a tool to record all the steps you will subsequently take to load, compile, initialize and solve the model. This tool will construct a script from the steps it sees you taking. You will likely then edit this script, for example to delete some of the missteps you have taken, and then save it. You may also pick out any of the steps in the script and execute them at any time rather than look for the tools in the tool set windows. You would use a script to aid you to repeat all the above steps quickly while you are debugging a model. You will also prepare a script to hand your model to someone else to execute. Indeed, your first experience with ASCEND may be to run a script that someone else has provided so you can be sure to run your first example successfully.



# CHAPTER 2 GETTING STARTED WITH ASCEND

## 2.1 PHILOSOPHY

Our goal is to create a set of very powerful modeling/solving tools. A side effect is that users can often find uses for the tools we did not anticipate. Another is that, while we have tried to build a user interface that lets every user from beginner to expert use our tools (or theirs combined with ours!) in a comfortable fashion, we have almost certainly erred on the side of giving the user too much control. Knowing that to be the case, the user should fearlessly dive in and try to use the system, at first by doing some of the simple problems we have provided in this documentation. The first step, of course, is to start the system up, the purpose of this section.

### 2.1.1 GETTING THE ASCEND SYSTEM AND INSTALLING IT

ASCEND is available through our Web page. Using your web browser, go the URL

`http://www.cs.cmu.edu/~ascend/Home.html`

Follow the instructions (the ftp link) there to download ASCEND.

#### Installing the UNIX version

If you are downloading a version to run on a UNIX workstation, then find someone who is a UNIX expert to help you. The process will involve transferring the source files for ASCEND along with a MAKE file. The MAKE file will allow a UNIX specialist to compile ASCEND and get it ready for use. There are detailed instructions that come with this version to help in installing it. (Your expert's expertise may be very minimally required for installing it on most systems.)

Installing the PC version

If you are downloading to a PC running under either NT or Windows 95, you will be downloading ASCEND4.zip. Uncompress using WinZip, double click on install.exe and follow the instructions.

**2.1.2 STARTING ASCEND****2.1.2.1 FOR PC USERS ONLY**

On the PC, simply double click on the ASCEND icon.

**2.1.2.2 FOR UNIX USERS ONLY**

The ASCEND IV interface is an open system written in TCL on top of several libraries of C code. The users are expected to customize it to suit their individual tastes.

We assume users are at least aware of the existence of environment variables and X resources. If you are not, contact your UNIX expert or the person who installed ASCEND on your system.

Environment Variables

Normally, if you are running on **UNIX** your system administrator will have set up a shell script to let you run ASCEND simply by typing `ascend`. To see if this is true, try typing

```
ascend -h
```

If this doesn't work, you may need to define the following environment variables in your `.login` (or perhaps `.profile`) file, or if you can find the ASCEND binary, it will frequently run without requiring a shell script.

**ASCENDDIST**

points to the directory where the ASCEND code has been installed.

```
setenv ASCENDDIST /usr1/ballan/asc4/test
```

**ASCENDHELP**

points to the ASCEND help file tree on your system. The tree does not have to reside with the rest of the distribution, though it may. This should have been configured for you when was installed.

**ASCENDLIBRARY**

is a colon-separated list of directories where ASCEND looks for files which are required by other files or which are read into ASCEND from a script without giving a complete path name. If you do not define ASCENDLIBRARY, the system will make guesses that usually work.

```
setenv ASCENDLIBRARY $ASCENDDIST/models/examples:$ASCENDDIST/models/libraries
```

## CHAPTER 3 SCRIPT

The Script Utility (see Figure 3-1) allows us to record the process of solving a model, or any other user interface process. Once this process is recorded in the form of a script, the script can be repeated either fully or in part. The solution process for a given model can be communicated to another modeler by distributing a script saved to a file. Following is an outline of the various menus and buttons on the script window along with a library of the ASCEND commands which can be recorded.

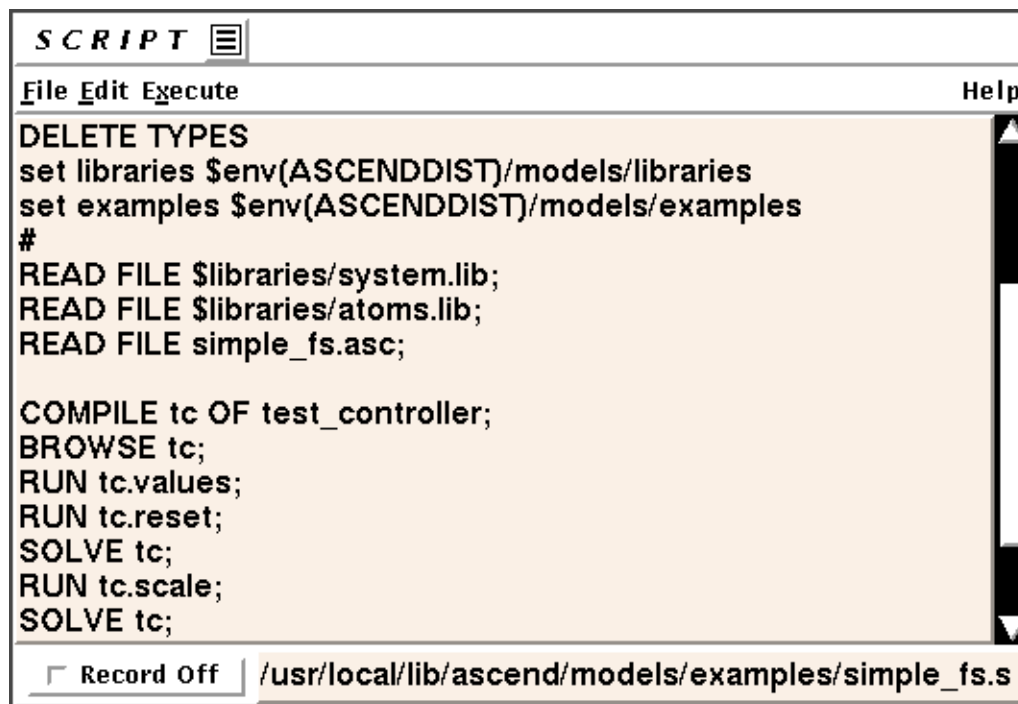


Figure 3-1 ASCEND's Script Window.

## 3.1 THE SCRIPT MENU BAR

### 3.1.1 SCRIPT FILE MENU

The script file menu provides several functions for managing script files. The script utility may contain multiple scripts but will only display one at any given time. Upon startup a scratch workspace is provided

<b>New File</b>	Request a buffer name and creates a new buffer with this name.
<b>Read File</b>	Requests a filename through the file selection box and proceeds to load this file (which is assumed to contain ASCEND Script and/or Tcl statements) into a new Script window buffer. No error checking is performed on the loaded file.
<b>Import File</b>	Requests a filename through the file selection box and appends the text contained in this file to the end of the current buffer.
<b>Save</b>	Saves the text in the current script buffer window to the current script file (indicated by the filename at the bottom of the script window). The existing file is overwritten.
<b>Save As</b>	Request a filename through the file selection box and saves the text in the current script buffer window to this file. If the specified file exists, it is overwritten.
<b>Buffer List</b>	A list of scripts used in the current ASCEND session is displayed at the bottom of the file menu. A script can be redisplayed in the script window by selecting it from the buffer list.

### 3.1.2 SCRIPT EDIT MENU

<b>Write selection</b>	Saves the <i>selected</i> (highlighted) text to a file which is specified through the file selection dialog box. Selecting text is discussed in Section 3.4.
<b>Select all</b>	Selects (highlights) all text in the window.
<b>Remove statements</b>	Removes (cuts) the <i>selected</i> text. The removed text is NOT saved for later pasting.

### 3.1.3 SCRIPT EXECUTE MENU

#### Statements selected

This button takes the selected text, breaks it into statements delimited by any semicolons (;) that appear in the selection, and executes each statement in the Tcl global environment.

### 3.1.4 SCRIPT TOOLBOX MENU

This menu has exactly the same content as the ASCEND toolbox window. See the chapter corresponding to the toolbox for details.

### 3.1.5 SCRIPT HELP MENU

#### On ASCEND/TCL Scripts

The highlighted selection is executed when you click Script.Execute.Selected\_statements. The selection is broken into statements at semicolons. Each statement is sent to Tcl for execution. If the statement doesn't return any errors, it is unhighlighted and the next statement is processed. The # character is the Tcl comment. It is most often used at the beginning of a line, and its effect stops at the end of a line. The SCRIPT automatically wraps lines too long for the window.

#### On SCRIPT

## 3.2 SCRIPT GRILL MENU

#### Record actions

When the record function is activated a log of interface events with defined ASCEND Script commands is appended to the end of the current script window buffer. Most, but not all, interface events have corresponding script commands. The record function can be turned on and off by toggling the pull down button on the grill menu or the record button at the bottom of the script window.

## 3.3 THE SCRIPT LANGUAGE

### 3.3.1 SUMMARY

Script keywords are commands defined for ASCEND (in CAPS) which may be used on the commandline or in the Script. Keywords are actually Tcl functions which encapsulate one or more of the C



primitives and other Tcl procedures, so that the user can conveniently emulate button presses. A working knowledge of tcl is not necessary to benefit from the Script's functionality; however, the tcl literate user will be able to create very powerful scripts.

Each keyword takes 0 or more arguments. The use of arguments is given in the following syntax:

<b>&lt;arg&gt;</b>	indicates the use of <b>arg</b> is required.
<b>&lt;a1 , a2&gt;</b>	indicates that the use of either <b>a1</b> or <b>a2</b> is required
<b>&lt;a1 a2&gt;</b>	indicates use of both <b>a1</b> and <b>a2</b> required. Usually written <a1> <a2>
<b>[a1]</b>	indicates the use of <b>a1</b> is optional.
<b>[a,b]</b>	indicates that either <b>a</b> or <b>b</b> is optional, but not both.
<b>qlfdid</b>	is short for 'QuaLiFieD IDentifier'
<b>qlfpid</b>	is short for 'QuaLiFied Procedure IDentifier'
	OF, WITH, TO, and other args in all CAPS are modifiers to the keyword which make it do different things.
<b>{ }</b>	It is generally best to <b>enclose all object names and units</b> in {braces} to prevent Tcl from performing string substitution.

### 3.3.2 QUICK REFERENCE:

<b>ASSIGN</b>	Sets the value of something atomic
<b>BROWSE</b>	Exports an object to the browser
<b>CLEAR_VARS</b>	Sets all the fixed flags to FALSE
<b>COMPILE</b>	Compiles a simulation of a given type
<b>DELETE</b>	Deletes a simulation or the type library
<b>DISPLAY*</b>	Displays something
<b>INTEGRATE</b>	Runs an IVP integrator
<b>MERGE</b>	Performs an ARE_THE_SAME

<b>PLOT</b>	Creates a plot file
<b>PRINT</b>	Prints one of the printable windows
<b>PROBE</b>	Exports an object to the probe
<b>READ</b>	Reads in a model, script, or values file.
<b>REFINE</b>	Performs an IS_REFINED_TO
<b>RESTORE*</b>	Reads a simulation from disk.
<b>RESUME</b>	Resumes compiling a simulation
<b>RUN</b>	Runs a procedure
<b>SAVE*</b>	Writes a simulation to disk
<b>SHOW</b>	Calls a unix plot program on a file from PLOT
<b>SOLVE</b>	Runs the solver
<b>WRITE</b>	Writes values in Tcl format to disk

\* Items not yet implemented.

### 3.3.3 COMMANDS

<b>ASSIGN</b>	ASSIGN <qlfdid> <value> [units]
	Sets the value of atom 'qlfdid' from the script. If value is real, it is required to give a set of units compatible with the dimensions of the variable. If the variable has no dimensions yet, ASSIGN will fix the dimensions.
<b>BROWSE</b>	BROWSE <qlfdid>
	Exports qlfdid to the browser, displaying it as the current instance in the browser.
<b>CLEAR_VARS</b>	CLEAR_VARS <qlfdid>
	Sets the value of the fixed flag to FALSE for all the variables on qlfdid.
<b>COMPILE</b>	COMPILE <simname> [OF] <type>

Build a simulation of the type given with name simname. You can get away with leaving out OF or spelling it wrong.

**DELETE** `DELETE <TYPES,simname>`

The modifier TYPES will cause all simulations to be deleted. If a simulation name (simname) is specified only that simulation will be deleted.

**DISPLAY** `DISPLAY <kind> [OF] <qlfdid>`

How qlfdid is displayed varies with kind. kinds are: VALUE  
ATTRIBUTES CODE ANCESTRY

**INTEGRATE** `INTEGRATE {qlfdid args}`

Runs an integrator on qlfdid. There are several permutations on the syntax. It is best to have solved qlfdid before hand to have good initial values.

```
INTEGRATE qlfdid (assumes LSODE and entire range)
INTEGRATE qlfdid WITH (assumes entire range)
INTEGRATE qlfdid FROM n1 TO n2 (assumes lode)
INTEGRATE qlfdid FROM n1 TO n2 WITH integrator
```

Requires:

- $n1 < n2$
- qlfdid be of an integrable type (a refinement of ivp.)

**MERGE** `MERGE <qlfdid1> [WITH] <qlfdid2>`

ARE\_THE\_SAME qlfdid1 and qlfdid2 if possible.

**OBJECTIVE** `OBJECTIVE`

Semantics of OBJECTIVE that will be supported are unclear as no OBJECTIVE other than the declarative one is yet supported. Not implemented yet

**PLOT** `PLOT <qlfdid> [filename]`

Writes plot data from qlfdid, which must be a plottable instance, to filename.

**PRINT** PRINT <PROBE,DISPLAY>

Prints out the text currently in the Probe or Display.

**PROBE** PROBE ONE qlfdid

Exports the item qlfdid to the Probe.

PROBE ALL qlfdid  
PROBE qlfdid

Exports items found in qlfdid matching the current specifications of Visit in the Browser. By default, all variables and relations.

Items always go to currently selected probe context.

**READ** READ [FILE,<VALUES,SCRIPT>] <filename>

Loads a file from disk. Searches for files in directories (Working directory):::\$ASCENDLIBRARY unless a full path name is given for filename.

The modifier FILE is used to indicate that the file contains ASCEND source code (ASCEND source code files normally have a .asc extension).

The modifier VALUES is used to indicate that the file contains variable data written by WRITE VALUES (These files normally have a .values extension).

The modifier SCRIPT is used to indicate that the file is a script file to be loaded at the end of the Script window (Script files normally have a .s extension).

If neither VALUES nor SCRIPT are found, FILE will be assumed.  
Note: You will get quite a spew from the parser if you leave out the SCRIPT or VALUES modifier by accident.

**REFINE** REFINE <qlfdid> [TO] <type>

Refines qlfdid to given type if they are conformable.

**RESTORE** RESTORE <file>

Reloads a simulation from disk

**RESUME** RESUME <simname>

Reinvokes compiler on simname.

**RUN** RUN <qlfpid>

Run the procedure qlfpid as if from the browser Initialize button.

**SAVE** SAVE <sim> [TO] <filename>

Filename will be assumed to be in Working directory (on utils page) unless it starts with a / or a ~. Not implemented yet.

**SHOW** SHOW <filename, LAST>

Invokes the plotter program on the filename given or on the file LAST generated by PLOT.

**SOLVE** SOLVE <qlfdid> [WITH] [solvername]

Exports qlfdid to the solver and attempts to solve it with the default solver (usually QRSlv) or the solver indicated by the optional solvername argument. Solvername must be given as it appears on the menu buttons. Bugs: Should use current solver rather than default.

**WRITE** WRITE <kind> <qlfdid> <file> [args]

Write something (what sort of write indicated by kind) about qlfdid to a file. args may modify as determined by kind. At present only VALUES is supported. SYSTEM (for solver dump) would be nice.

WRITE VALUES filename.

Filename must be a full path name or in the pwd, also known as ‘.’.

## 3.4 SCRIPT WINDOW BINDINGS

In the event binding descriptions that follow, M1 is short for mouse-button-1 (the left mousebutton), M2 is the middle button, and M3 is the right mouse button. On machines with no middle button, M3 is still the right mouse-button and M2 is unavailable.

<b>M1</b>	repositions the cursor.
<b>M1-Drag</b>	selects text.
<b>Shift-M1 [-Drag]</b>	extends the selection.
<b>Double-M1</b>	selects the nearest word.
<b>Double-M1-Drag</b>	selects the nearest word and those you drag over, whole words at a time.
<b>Triple-M1</b>	selects the nearest line.
<b>Triple-M1-Drag</b>	selects the nearest line and those you drag over, whole lines at a time.
<b>M2</b>	does nothing.
<b>M2-Held-Down</b>	has an effect similar to the scrollbar.
<b>M3</b>	does nothing.
<b>Control-M1</b>	Starts another part of a disjoint selection.
<u>UNIX bindings:</u>	The text widgets in ASCEND share a common stack of cut/copy/paste text pieces. This is a CMU extension of the text bindings, not default Tk behavior, and it is EMACS-like, but not EMACS (EMACS uses a ring, not a stack.) When the stack is empty, Paste does nothing. This is a design decision. The Tcl function ascPopText can be changed to behave differently.
<b>Control-k</b>	Cuts text to the end of the current line, putting it on the stack.
<b>Control-w</b>	Cuts the selected text, putting it on the stack.
<b>Meta-w</b>	(e.g. diamond-w on most Sun keyboards) Copies the selected text onto the stack.
<b>Control-y</b>	Pastes the most recent text added to the stack, and removes it from the stack.
<b>Meta-y</b>	Not supported.
<u>MSW bindings:</u>	The standard Control-X, Control-C, Control-V bindings of Microsoft Windows clipboard apply to text widgets. The UNIX text stack is not available.



## CHAPTER 4 LIBRARY

The Library window (Figure 4-1) in ASCEND allows the user to read *types* into the ASCEND system from files, *compile* types into instances, and delete types.

Types are the templates used to create simulations. They come in two flavors: ATOM, which has a value associated with the instance name when it is instantiated, and MODEL, which has no value. ATOMs, further, come in vanilla and UNIVERSAL flavors. Universal atoms have a single compiled instance which is global to all simulations created.

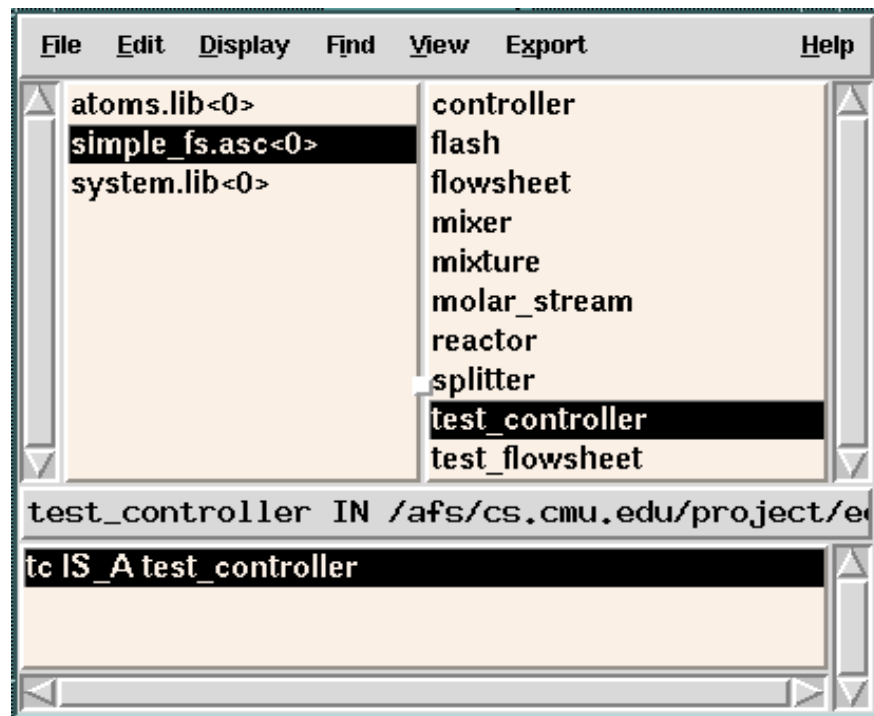


Figure 4-1 ASCEND Library Window.

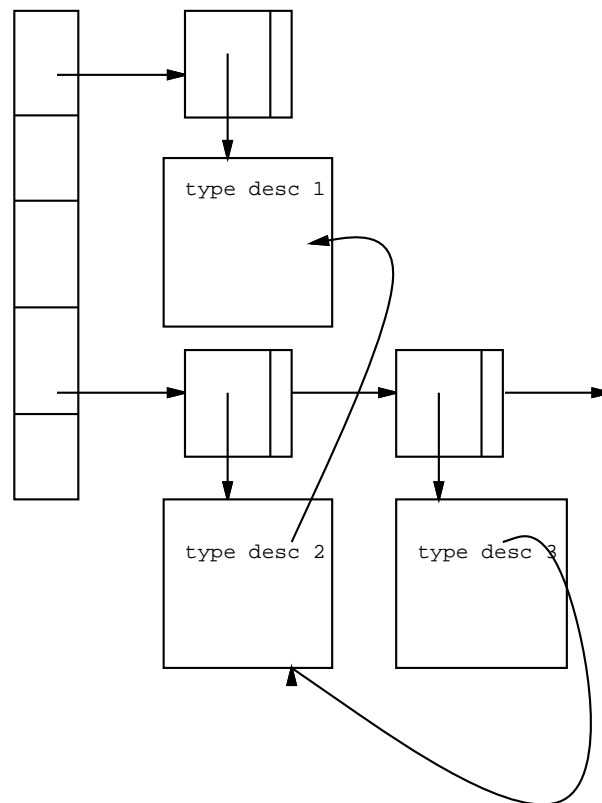


Both ATOMS and MODELS are defined in source files. By convention, source files are named with the endings `.asc` and `.lib`, though other extensions may be used.

In the ASCEND Library window, source files appear in the upper left box. On the other hand, the types defined in the highlighted source file appear in the upper right box. A double-button2 in either box will compile the highlighted type definition. It doesn't reselect. The upper left box should perhaps have double-button2 bound to reread the selected source module. The ASCEND fundamental type such as integer, real, etc., are not shown in the library window, since their definition is performed internally, not by using a specific source file. The lower box of the ASCEND Library window contains the name of the simulations that have been compiled and can be run.

The data structure used to store type definitions is sketched in Figure 4-2.

Type Library



Notes: type desc3 has a refinement ptr to type desc 2  
 type desc2 has a refinement ptr to type desc 1

The problem is when type desc 2 is being redefined  
 by reloading a new module.

Figure 4-2 Data structure used to store type definitions.

## 4.1 MENU BAR

The menu bar on the Library window has seven entries: File, Edit, Display, Find, View, Export and Help.

### 4.1.1 THE FILE MENU

#### **Read types from file**

This loads type definitions into the system. The file selection dialog is used to select a source file.

The names of types are unique within the system. A new definition of a type overwrites the old definition of a type in all cases. If the new definition and the old definition were read from files of the same name, this overwrite will be done silently. If the new definition comes from a different file, the overwrite will be done noisily.

**This is incorrect, but perhaps is as it ought to be.** *Existing types which refined or had parts that were of the old type definition will now refine or have parts which are of the new type. e.g. If you reread system.asc (and hence solver\_var) everybody in the interface library who pointed at the old solver\_var type will now point at the new solver\_var type.*

Instances already compiled using the definitions that have been overwritten will continue to point at a copy of the old definition the system has squirreled away somewhere. These squirreled away copies will not necessarily be the same as what is in the interface type library if you have reread a file with a newer type definition. This may cause refinement of the old instance to fail. In general if you redefine a type, you will probably want to re instantiate things that depend on that type.

#### **Close window**

It just closes the ASCEND's Library window.

### 4.1.2 THE EDIT MENU

#### **Create simulation**

Create (or instantiate) a simulation based on a type definition. Anytime that the compile button is selected, the compile dialog window shown in Figure 4-3 will ask for the name which will be used to identify the simulation. All simulation created can be seen and in the lower box of the ASCEND Library window. This box can contain any number of simulations.

There is a second way to create a simulation through the *Script* window. This option is explained in the ASCEND Script window document.

**Delete Simulation**

Once a simulation has been loaded into the lower box of the Library window, it can easily be removed by selecting this option from the **Edit** menu.

**Delete all types**

Destroys all simulations and deletes all types. This option has no effect in the fundamental definitions.

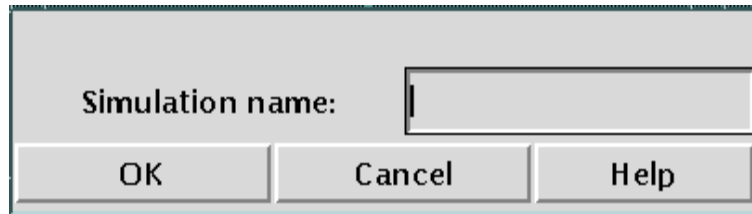


Figure 4-3 The Create Simulation Dialog

### 4.1.3 THE DISPLAY MENU

Most of the options in the Display Menu will be enabled only if a type definition has been selected, this is because the tasks performed in the menu are implicitly associated with a type definition.

**Code**

Displays the source code of the selected type in the ASCEND Display Window.

**Ancestry**

Allows the use of the Type Refinement Hierarchy Window. See the section corresponding to this topic.

**Refinement hierarchy**

Displays the refinement hierarchy of the selected type in the ASCEND Display window.

**External Functions**

It reports any external function defined from a loaded package library. The list of external functions (if any) is displayed in the window from which ASCEND IV was started

**Hide Type**

Any type definition which is selected to be hidden will be ignored for browsing purposes. Internally, the selection of this option consists in setting to zero a binary flag included in the type description of the highlighted type definition.

**UnHide Type**

Reverses the action of “hiding” a type. As a consequence, such a type will be considered for browsing purposes. Hide Type and UnHide Type are never enabled at the same time (for the same type definition)

because they have opposite meaning. The default for all the type definitions (except fundamentals) is to be “unhidden”.

Both Hide Type and UnHide Type have two selections as a submenu. The user can ask for the un/hiding of only the **selected type**, or for the un/hiding of the selected **type and its refinements**.

### Hide/Show Fundamentals

This special option is given because fundamental types do not appear as definitions in the ascend libraries, but we still may want to able/enable such types for browsing purposes. When this button is selected, the window shown in Figure 4-4 will be used to perform the desired hiding or un hiding of any of the fundamental types.

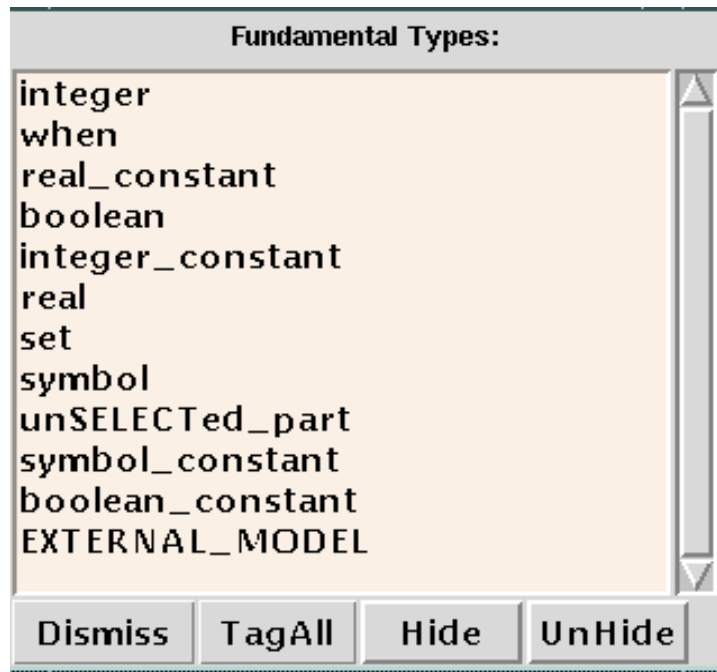


Figure 4-4 Select the fundamental type to Hide or Unhide.

## 4.1.4 THE FIND MENU

### Type by name

Finds a type by its name.

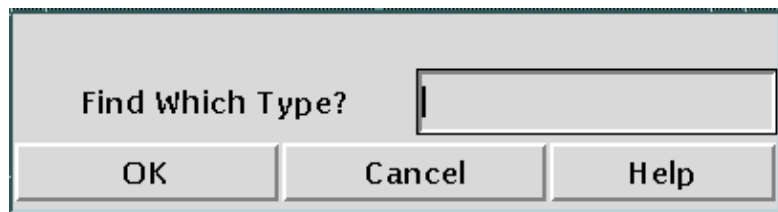


Figure 4-5 The Library's Find Type dialog.

<b>Type by fuzzy name</b>	Finds all type names that match a word (provided by the user) in any fuzzy way. For instance, the name column could give as a result the following list: demo_column, mw_demo_column, plot_column, etc. The fuzzy name is defined in a dialog window similar to that used in the Find Type by name option.
<b>Pending statements</b>	There are three selections under the Pending Statements submenu, these are <b>To Display</b> , <b>To Console</b> , and <b>To File</b> . Pending in a simulation are relations that have not yet been fully processed by ASCEND's compiler. It is the modeler's job to correct the pending relations in order to arrive at a fully functional simulation. Corrections may be made by either creating a model which refines the current model or by editing ASCEND code and starting over. This option gives the user access to information about the type and location of the pending statements.
<b>To Display</b>	By selecting the <b>To Display</b> option, all of the simulation pendings are displayed in the <i>Display</i> window.
<b>To Console</b>	By selecting the <b>To Console</b> option, all of the simulation pendings are displayed in the window from which ASCEND IV was started.
<b>To File</b>	By selecting the <b>To File</b> option, the <i>File select box</i> is opened and the user is asked to enter the name of the file in which to save the model pendings.

#### 4.1.5 THE VIEW MENU

This option has the same application in all of the ASCEND windows and is explained as a general tool in a companion document.

#### 4.1.6 THE EXPORT MENU

There are three selections under this submenu, these are **Simulation to Browser**, **Simulation to Solver**, and **Simulation to Probe**.

<b>Simulation to Browser</b>	By selecting the <b>Simulation to Browser</b> option, the simulation highlighted in the lower box of the Library window is loaded into the <i>Browser</i> . From the <i>Browser</i> , the model can be explored in more detail.
<b>Simulation to Solver</b>	By selecting the <b>Simulation to Solver</b> option, the simulation highlighted in the lower box of the Library window is loaded into the <i>Solver</i> . (Note that exporting to the solver causes a degrees of freedom analysis to be carried out.)

**Simulation to Probe** By selecting the **Simulation to Probe** option, all of the variables of the simulation highlighted in the lower box of the Library window are loaded into the *Probe*. This is not recommended as there are usually more variables in a model than the user would wish to view at one time. However, if the user does wish to look at all of the variables and their current values, the **Simulation to Probe** option can be useful.

#### 4.1.7 THE HELP MENU

**On LIBRARY** Brings up a browser pointing to the information provided in this document.

## 4.2 TYPE REFINEMENT HIERARCHY WINDOW

The type tree is a directed acyclic graph (DAG) based on the type hierarchy currently defined in the interface Library. Selection of the option Display Ancestry for any selected type gives the entire refinement hierarchy for that type, by enabling the use of the window shown in Figure 4-6.

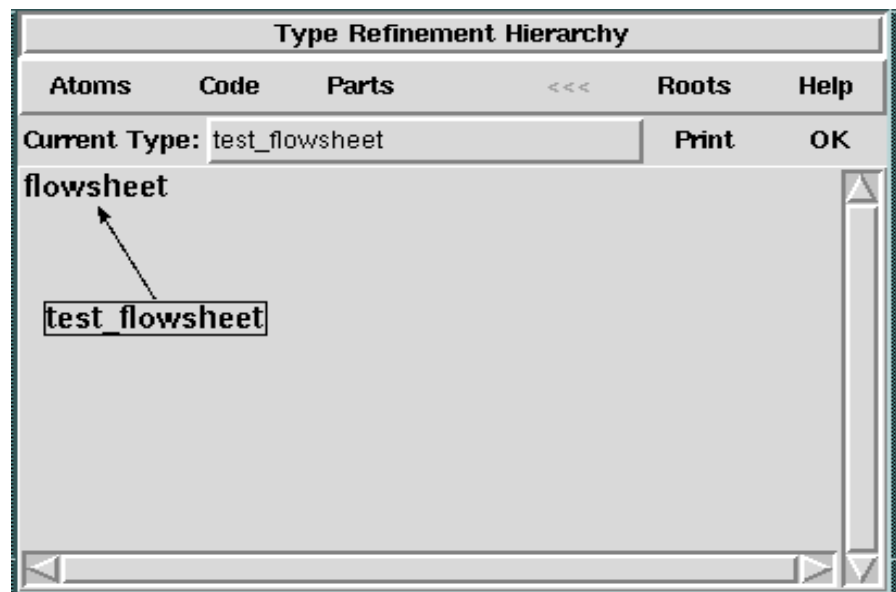


Figure 4-6 The Type Refinement Window.

The current focus in the hierarchy is indicated by a rectangle around the type name and the Current type.

The buttons on the left in the type window operate on the currently selected type:

'Atoms' shows the types of ATOMIC parts in the selected type definition. It also shows the incremental code for the type. You can select from the part types list to look at a different hierarchy.

'Code' shows the internally stored code of the selected type. The expressions, both algebraic and logical, are in reverse Polish notation. This is different from the way the code of the Library Display Code button shows it. Comparison of the two is sometimes a useful debugging tool.

'Parts' (Figure 4-7) shows the types of MODEL parts in the selected

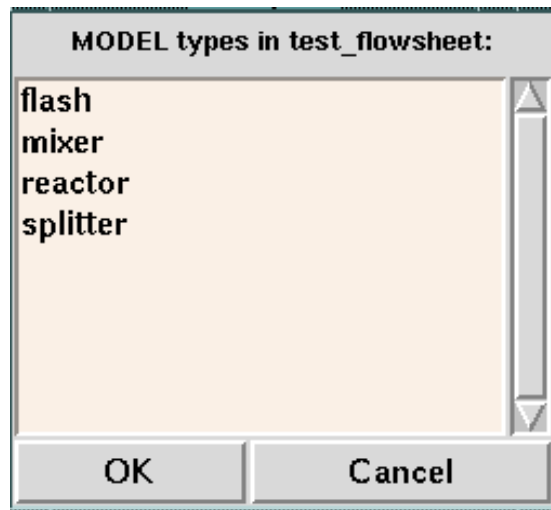


Figure 4-7

The Parts window displays the parts.

type definition. It also shows the incremental code for the type.

The '<<<' (or backtrack) button backs up to the previously displayed type hierarchy, if there is one.

'Roots' (Figure 4-8) shows the existing root types, that is, the existing types which are not refinements of anything.

While ASCEND is building the graph, you may see a spew in the window from which ASCEND was started about orphaned types. This means there are types in the Library which are refinements of older types which are no longer in the Library.

While ASCEND is getting the Atom or Model parts list for a type, part types names which are undefined will be spewed.

When an older type is replaced in the Library by a new one of the same name, the old one is squirreled away where types that refined it can still see it. The only way to get current types to look at the new definition without touching the source files for the current types is to delete all types and reread the entire Library.

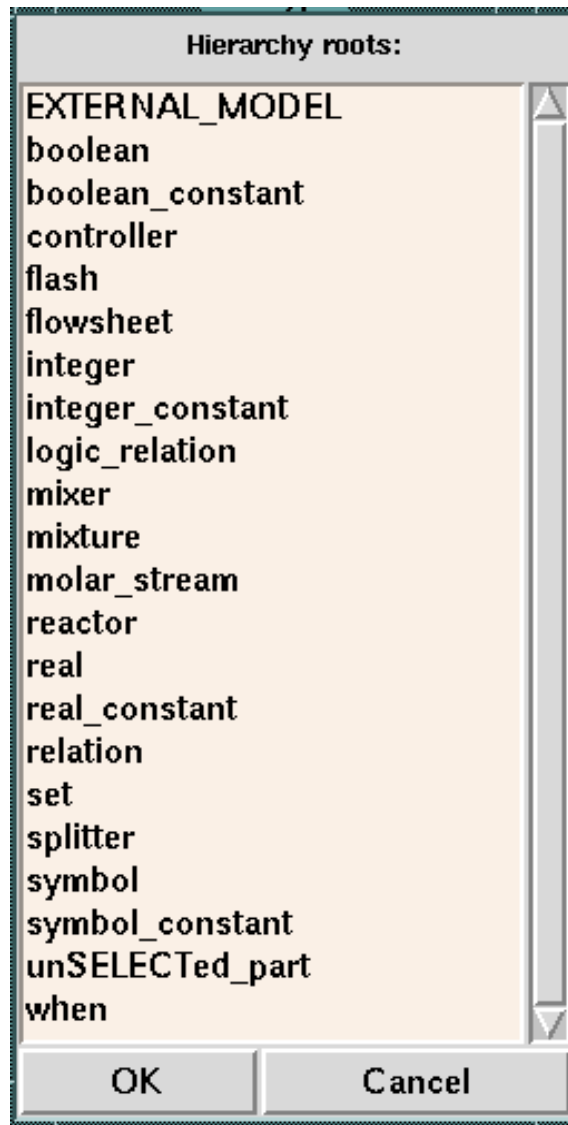


Figure 4-8 The Hierarchy Roots Window.





# CHAPTER 5 MERGED INTO LIBRARY

## 5.1 SIMS WINDOW

The *Sims* window, short for Simulations window, contains the name of the simulations that can be run. We can see in Figure that there are three major menus and the **Help** menu available through the Sims window.

Figure 5-1



The menus are the **Edit**, **Pendings**, and **Export** menus. There are two ways a simulation can be created. The first way is manually through the Library window. Assume for the moment that you are running a reactor model and the following libraries and models have been loaded into the *Library* window.

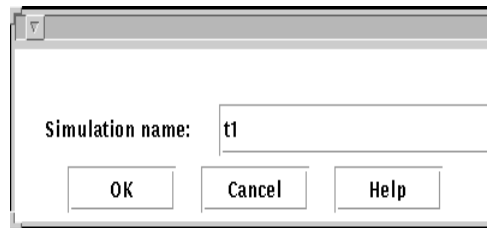
```
ivpsystem.lib;  
atoms.lib;  
components.lib;  
H_G_thermodynamics.lib;  
stream.lib;  
reactor.lib;  
test_reactor.asc;
```

By highlighting test\_reacotr.asc, the following options appear in the right hand side section of the *Library* window.

```
reactor_control
reactor_test
td_reactor_test
```

If you highlight `reactor_test` option and select **Compile** from the **Create** menu, a window appears that allows the user to enter the name of the simulation. In this case, we will simply call the simulation `t1`. Figure shows this window.

Figure 5-2



Once the simulation name has been entered, select the OK button. Once this is complete, we see that

```
t1 IS_A reactor_test
```

has been added to the *Sims* window. The *Sims* window can contain any number of simulations.

The second way to create a simulation is through the *Script* window. Assuming that the necessary libraries and models have been loaded into the *Library* window, simulation `t1` above can be created by executing the following line in the *Script* window.

```
set model t1;
```

We see that the second option is not only faster, but it is more convenient.

## 5.1.1 THE EDIT MENU

There are three selections available in the **Edit** menu, these are **Delete**, **Save**, and **Restore**.

### 5.1.1.1 DELETE

Once a simulation has been loaded into the *Sims* window, it can easily be removed by selecting the **Delete** option from the **Edit** menu.

### 5.1.1.2 SAVE

This option is currently not functional.

### 5.1.1.3 RESTORE

This option is currently not functional.

## 5.1.2 THE PENDINGS MENU

There are three selections under the **Pendings** menu, these are **To Screen**, **To Display**, and **To File**. Pendings in a simulation are relations that have not yet been fully processed by ASCEND's compiler. It is the modeler's job to correct the pending relations in order to arrive at a fully functional simulation. Corrections may be made by either creating a model which refines the current model or by editing ASCEND code and starting over. The Pendings menu gives the user access to information about the type and location of the pending statements.

### 5.1.2.1 TO SCREEN

By selecting the **To Screen** option from the **Pendings** menu, all of the simulation pendings are displayed in the window from which ASCEND IV was started.

### 5.1.2.2 TO DISPLAY

By selecting the **To Display** option from the **Pendings** menu, all of the simulation pendings are displayed in the *Display* window.

### 5.1.2.3 TO FILE

By selecting the **To File** option from the **Pendings** menu, the *File select box* is opened and the user is asked to enter the name of the file in which to save the model pendings.

## 5.1.3 THE EXPORT MENU

There are three selections under the **Export** menu, these are **to Browser**, **to Solver**, and **to Probe**.

### 5.1.3.1 TO BROWSER

By selecting the **to Browser** option from the **Export** menu, the simulation is loaded into the *Browser*. From the *Browser*, the model can be explored in more detail. This is covered more thoroughly in the *Browser* section of the documentation.

### 5.1.3.2 TO SOLVER

By selecting the **to Solver** option from the **Export** menu, the simulation is loaded into the *Solver*. (Note that exporting to the solver causes a degrees of freedom analysis to be carried out.)

### 5.1.3.3 TO PROBE

By selecting the **to Probe** option from the **Export** menu, all of the variables of the simulation are loaded into the *Probe*. This is not recommended as there are usually more variables in a model than the user would wish to view at one time. However, if the user does wish to look at all of the variables and their current values, the **to Probe** option can be useful. More on this is covered in the *Probe* section of the documentation.

## CHAPTER 6 BROWSER

The Browser window (Figure 6-1) provides the means with which to view the parts of a simulation. When a simulation is exported to the Browser, the name of the simulation appears in the Browser's upper left box and the child instances of the simulation appear in the upper right box. Selecting a child instance in the right box will move the instance to the bottom of the stack in the left box and display its children in the right box. The instance tree can be traversed in this manner until an atom (usually a variable) resides at the bottom of the stack in the left box and its attributes appear in the right box. Selecting a member of the stack in the left box will clear any lower instances on the stack and display the selected instance's children in the right box.

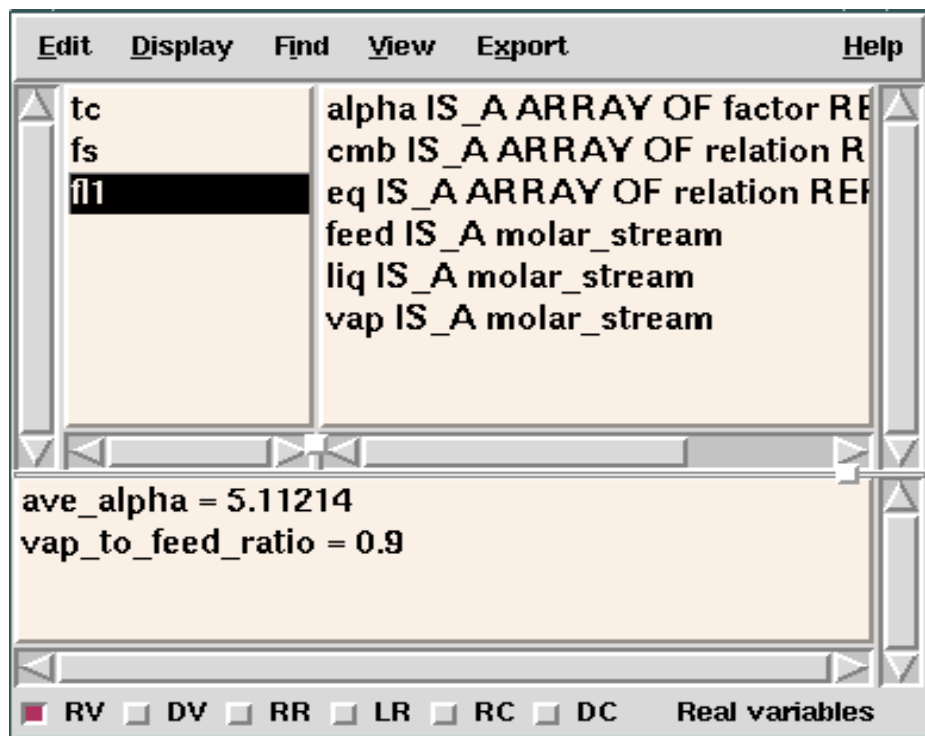


Figure 6-1 ASCEND's Browser window.

A subset of the instances appearing in the upper right box as well as the values of these instances appear in the Browser's lower box. Which subset of instances appears in the lower box is controlled by the user by clicking in some of the options given in the bar at the bottom of the Browser window. In Figure 6-1 RV has been selected. RV stands for **Real Variables**. Therefore, the child instances of fl1 which are real variables and the values of these real instances are shown in the lower box. Other options in the bar at the bottom of the Browser window, which can be simultaneously selected, are DV (Discrete Variable), RR (Real Relations), LR (Logical Relations), RC (Real Constants) and DC (Discrete Constants). Selecting an instance in the lower box with the right button of the mouse will have the same effect as selecting the same instance in the upper right box. On the other hand, selecting an instance in the lower box with the left button of the mouse will bring up the "Set Value" Dialog box, which will give the user the option of modifying the value of the selected instance. More about the "Set Value" option will be given in the following section of this document.

## 6.1 THE MENU BAR

The menu bar on the Browser window has six entries: Edit, Display, Find, View, Export, and Help.

### 6.1.1 BROWSER EDIT MENU

#### **Run method**

If the instance in the left box has one or more methods available, Edit ->Run Method will be available for selection. Selecting Run Method will display the Methods Selection Window containing a list of available methods for the current Browser instance. A method is selected by clicking its name (only one method can be selected at a time). Depressing the OK button will run the selected method. Depressing the Show button will display the code for the selected method. Depressing the Cancel button will close the Method Selection Window without running any method.

#### **Clear Vars**

In ASCEND, the type solver\_var and all its refinements constitute a variable for solution purposes. Each variable has a boolean, named "fixed", as one of its children. When a variable's fixed boolean, or fixed flag as it is commonly called, is set to False, the variable is considered an output variable (i.e. the solver will determine its value). The Clear Vars method sets the fixed flag of every variable which is a child of the current Browser instance to False.

#### **Set value**

When the current Browser instance is a real, symbol, integer, or boolean Edit->Set Value will be available for selection. Selecting Set Value displays the Set Value Dialog box. The value (and units in the case of reals) may be set by filing in the value (and units) fields of the Set Value Dialog box and depressing the OK button. Depressing the

Cancel button closes the Set Value Dialog box. Booleans are assigned simply by double clicking the mouse button 2 on their name when it appears in the right browser box. Write values

Selecting Edit->Write Values saves the attribute values of all atoms which are descendents of the current instance to a file. A file select box is displayed so a new file may be created or an old file over written. The attribute values are written to the selected file along with their path names relative to the current instance. The first line of the file specifies the path from the simulation to the current instance.

**Read values**

Reads the values from a file previously saved by Write Values. Values files are read using full path names (including the simulation name). The simulation for which values are being read does not necessarily have to be in the Browser (but it should exist). The first line a values file may be edited in order to read values to an instance with a path name other than the path indicated in the file.

**Refine**

Refines the current Browser instance to a given type. Edit->Refine may only be selected if the Library contains types which are refinements of the current Browser instance type. Selecting Edit->Refine displays the eligible types for the refinement of the current part in the Refinement dialog box. Selecting a type and depressing OK refines the current type to the selected type. Depressing Show displays the ASCEND code for the selected type. Depressing the Cancel button closes the Refinement dialog box without making any refinements.

**Merge**

ARE\_THE\_SAME the current part (left side of the Browser) with another given part. Do not ARE\_THE\_SAME parts from 2 different simulations. You cannot merge parts of atoms being atomic with anything. The dialog box will ask for the name of the instance that you want to merge with the instance highlighted in the left box of the browser.

**Compile**

Submenu containing Resume Compilation and Create Part.

**Resume  
Compilation**

Attempts to process any pending statements in the simulation in the Browser. It does not matter where in the simulation you have browsed to, Resume always starts at the top.

**Create Part**

This is a feature of the PASCAL version only. The proper way to add a part to a simulation is to create a refinement of the original model in a new file, read in that definition, and refine the simulation up to that new model.



## 6.1.2 BROWSER DISPLAY MENU

<b>Attributes</b>	Display the attributes of a real variable. Other functionality may be added later to this button.
<b>Relations</b>	Display all the relations at or below the current point in the Browser. Relations get arbitrary names unless explicitly named by the user in code. The arbitrary name, at the moment consists of ParentName_n where n is the number of the nth relation in the MODEL ParentName. If this name is not unique, enough letters a-z get added to make it unique. When the instance highlighted in the left box of the Browser is a real variable, this option will display all of the relation in which such a variable is incident.
<b>Cond Rels</b>	Display all the conditional relations at or below the current point in the Browser. Conditional Relations do not have to be satisfied. They are used as boundaries in conditional programming. The arbitrary name of a conditional relation is obtained in the same way as any other relation, but in general, the name of a conditional relation must be provided by the user, since the operator SATISFIED requires such a name.
<b>Log Rels</b>	Display all the logical relations at or below the current point in the Browser. Logical Relations get arbitrary names unless explicitly named by the user in code. The arbitrary name of a Logical Relation follows the same pattern as that of real relations. When the instance highlighted in the left box of the Browser is a boolean variable, this option will display all of the logical relation in which such a boolean variable is incident.
<b>Cond Log Rels</b>	Display all the logical relations at or below the current point in the Browser. Conditional Logical Relations do not have to be satisfied. They are used as conditions to check in conditional programming. The arbitrary name of a conditional logical relation is obtained in the same way as any other logical relation, but in general, the name of a conditional logical relation must be provided by the user, since the operator SATISFIED requires such a name.
<b>Whens</b>	This option is enabled for instances of models, relations, booleans, symbols, and integers. For the case of a model instance, this button will display not only all the when instances defined as parts of such a model, but also the when instances which include such a model in one of their CASEs. Distinction is made between those two possibilities. For relation, boolean, symbols and integer instances, this option displays the when instances which include such relation, symbol, etc., either in one of their CASEs or in the list of conditional variables.

When instances are useful for the conditional configuration of a problem and always get arbitrary names.

**Plot** Invokes a plotting program, if allowable, on the current object. The type of plot generated is controlled by the Utilities page variables Plot.type and Plot.program.

**Statistics** Prints out some information about the object tree in the Browser starting from the current point and going downward.

### 6.1.3 BROWSER FIND MENU

**By name** Search for an instance with a given qualified name and go there. The name of the instance to search for is defined in the dialog. This option may be useful for jumping around in the instance tree.

**By type** You can search for objects of any particular type with certain attributes. The allowable searches are best explained by examples as shown in Table 6-1. The search is loosely matched, i.e. any object that is of the

**Table 6-1** Examples of the performance of the Find by type option

Type	Attribute	Low Value	High Value	Explanation
unit				Find all parts that are units.
solver_var	fixed			Find all refinements of solver_var with a part fixed
solver_var	fixed	TRUE		Find all refinements of solver_var with a part fixed where fixed==TRUE
stream	Ftot	4		Find all streams with part Ftot where value is $4 \pm \text{epsilon}$
stream	Ftot	4	10	Find all streams with part Ftot where $4 \leq \text{Ftot} \leq 10$
relation	VALUE	0		Find all relations with a residual $0 \pm \text{epsilon}$
symbol	VALUE	ACH		Find all symbols where VALUE is 'ACH'
symbol	VALUE	ACH	ACZ	find all symbols where 'ACH' $\leq$ VALUE $\leq$ 'ACZ'

**Table 6-1** Examples of the performance of the Find by type option

Type	Attribute	Low Value	High Value	Explanation
component_constant				Find all parts that are component_constants
symbol_constant	VALUE	UNDEFINED		Find all undefined symbol_constants. Works for all types with a value.

type given, OR a refinement of the type given and matches the attribute qualifications, will be on the list of items found.

If there are no matches, there is no results box: just a message in the command line or a popup error message.

The results of the Find appear in a box and you can export 1 or more of the results in the box to the Browser or the Probe by selecting them and clicking on Browser or Probe. When you have finished exporting items to wherever you like, click on OK. The rest of the interface will ignore you until you dismiss the box.

Notes:

- Clear any of the extra fields not required for your search before you hit OK. We will usually find nothing that matches if there are extra search parameters hanging around that don't make sense.
- VALUE is a special keyword for dealing with atomic types. Variables and symbols have a value internally but not a child named VALUE. Similarly, relations have a residual but not as an accessible part at the moment.
- Symbols and integers will be matched exactly if only a low value is given. The matching of symbols given a low and high value is done lexically according to the collating sequence of the machine in use.
- Frequently what you really want to see is the name of a set of things of a given type. (E.g. number 8 where you want to know what components are in a flowsheet.) Find will return the instances though, not their common parent. Simply export one to the Browser and then click up a level to see the set of components in use.
- You can tab between fields in the Find by Type widget.
- You can select a type name in the library, call up the Find by Type search in the Browser, put the mouse cursor over the Type entry

and hit Ctrl-Y to copy name of the type selected in the library. This works on most window managers. If you get something other than the type name you expected, just hit Ctrl-U to empty the entry and type the name in yourself.

- Epsilon is about  $1e-8$  in terms of the SI units for any real quantity.

### **Aliases**

Find all the other names that the current object has in the simulation. For example, assume that you have named a simulation as fs. Assume further that the output stream from the mixer, m1, is merged with the input stream for the reactor, r1. Then, that stream is an object with two different names. Suppose you are looking at r1.feed as the current object. Asking for aliases will give the list

fs.r1.feed

fs.m1.output

If you pick one of the aliases, it can be exported to the BROWSER, the SOLVER or the PROBE. Alternate names for objects can also be created by ALIASES statements and by passing them into a parameterized MODEL, not only by merging.

### **Where created**

Find the other names that the current object was CONSTRUCTED under. If an object is shown as being created under 4 names, it means that once there were 4 objects and that 3 were destroyed in merge (ARE\_THE\_SAME) statements to reach the current unity. (Merging is expensive).

If you pick one of the names, it can be exported to the BROWSER, the SOLVER or the PROBE. Alternate names for objects can also be created by ALIASES statements or by passing them into a parameterized MODEL, but these names do not appear in the list of creations.

### **Clique**

Find all the instances that ARE\_ALIKE with the current one. The instances shown are bound together so that if the formal type of one is changed, they are all upgraded with the first. Parameterized objects cannot be ARE\_ALIKE'd because when parameters exist the formal type requires outside information (the parameters) in order to check that it is being used in a valid way.

### **Eligible variables**

Find real variables eligible to be fixed. If the solver is occupied by the same simulation, this query is thrown to the Solver. If not, the degrees of freedom are analyzed as if the current model were exported to the Solver.

<b>Relations</b>	Not implemented. See Export for ways to find relations and send to the Probe.
<b>Operands</b>	Not implemented.
<b>Parents</b>	Not implemented.
<b>Pendings</b>	Pendings has been moved to the Library window.

### 6.1.4 BROWSER VIEW MENU

The first three options in the View menu are toggles that determine your preferences when browsing instances:

<b>Suppress Atoms</b>	This button toggles whether or not to show atomic instances in the upper right box of the Browser window.
<b>Display Atom Values</b>	This button toggles whether to display values or to display the types of atoms in the child box (upper right side) of the Browser. For the case of relations, the residual shown with the relation is the last computed by the solver and not the residual at the current values of the variables.
<b>Check Dimensionality</b>	This switch turns warnings about relation inconsistency off and on. In principle it should not be necessary, but for the quick and dirty model it is sometimes handy.
<b>Hide Names</b>	This option has a similar functionality from that of Hide Types in the ASCEND Library windows. That is, it will hide or unhide instances for browsing purposes. The difference, however, is that this option hides by name, not by type. To clarify, it is quite different to hide instances of name fs from to hide instances of type test_flowsheet.
<b>UnHide Names</b>	Reverses the effect of the command Hide Names. By default, all the names are “unhidden”, therefore, this option is used only after some of the names has been hidden.

The final options in the View menu correspond to general features in the View menu of any of the ASCEND windows, and they are explained in a companion document.

### 6.1.5 BROWSER EXPORT MENU

<b>to Solver</b>	Checks the model for exportability (must be of type MODEL without any pending compilation) and sends it to the Solver.
------------------	--

**Many to Probe**

Sends the child instances of the current part being browsed to the Probe. The types of instances sent to the Probe are selected in the filtering window shown in Figure 6-2. Every switch toggles whether or

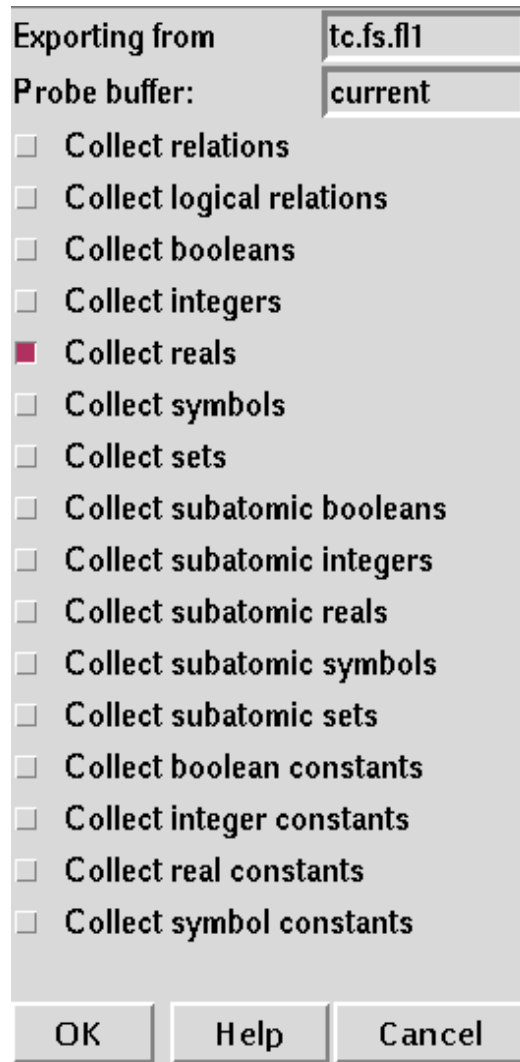


Figure 6-2 Filtering instances sent to the Probe

not to export each of types to the Probe.

**Item to Probe**

Exports the instance on the left box of the Browser to the Probe.

### 6.1.6 BROWSER HELP MENU

**On BROWSER**

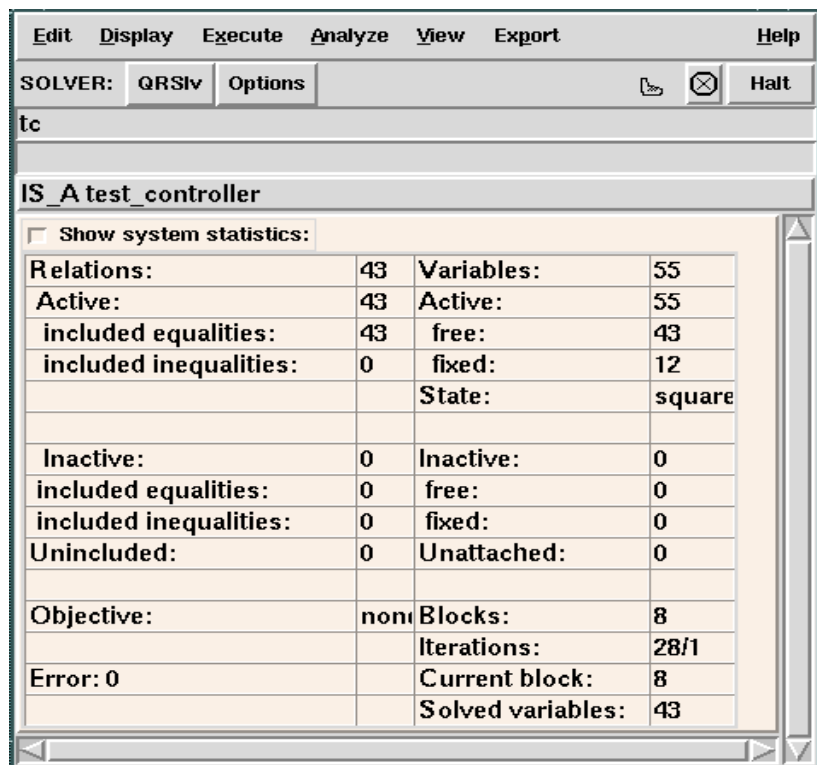
Brings up a browser pointing to the information provided in this document.



# CHAPTER 7 SOLVER

The purpose of the [Solver Utility](#) shown in [Figure 7-1](#) is to provide support for the numerical solving and debugging of an imported instance. To this end the Solver allows the user to access numerical solvers and analysis functions and displays statistical and status information for the problem being solved. The upper section of the solver window contains a menu bar; buttons for selecting numerical solvers, solver options, and halting the solver; a label containing the name of the current instance (or problem being solved); and a label containing the type of the current instance. The remainder of the Solver window is devoted to providing statistics about the problems relations and variables along with a description of the problem's state.

Figure 7-1 Solver Window





## 7.1 THE SOLVER MENU BAR

The menu bar on the Solver window has seven entries: Edit, Display, Execute, Analyze, View, Export, and Help.

### 7.1.1 SOLVER EDIT MENU

- Remove instance** Removes the current instance from the solver.
- Select objective** Provides a list of objectives from which one may select. The selected objective will be used in any subsequent optimizations until another objective is selected.

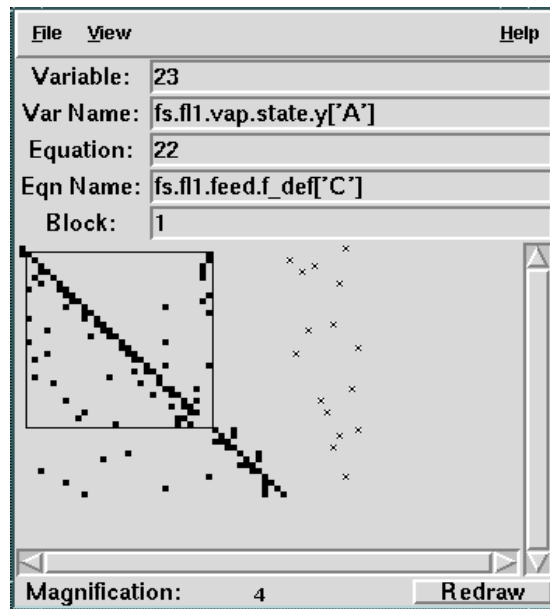
### 7.1.2 SOLVER DISPLAY MENU

- Status** Shows the internal status of the Solver along with the largest block scaled residual vector two-norm.



- Unattached variables** Shows variables not incident in any of the relations in the current system being solved.
- Unincluded relations** Shows relations not in the current system being solved.
- Incidence matrix** Incidence matrix shows the incidence of variables in relations (See Figure 7-2). Clicking mouse-1 (left button) on the matrix displays the names and numbers of the relation/variable at that coordinate, whether that coordinate is occupied or not. A box is drawn around the partitioned block containing the selected coordinate and the block number is displayed. The selected block or the entire incidence matrix may be printed by selecting the PrintBlock or the Print button respectively. The scale of the incidence matrix can be changed by sliding the magnification bar and depressing the Redraw button. Depressing the OK button will close the INCIDENCE window.
- Drawing large dense matrices may take a while. Drawing matrices on problems bigger than about 1000x1000 may be prohibitively expensive on slow machines. The row/column ordering is that done by the selected solver, except that fixed vars and unincluded relations are moved to the edges.

Figure 7-2 The Incidence Matrix



### 7.1.3 SOLVER EXECUTE MENU

- Solve** Solve the current problem as an algebraic or optimization problem depending on what solver is selected
- Single step** Perform a single iteration of the system with the solver in question. In some solvers (e.g. MINOS) there is no iteration mode. For these selecting single step will result in a full solve attempt.
- For QRSly, an iteration will be a Newton like step if there are many variables in the current block or if the current block is a blackbox singleton. Singletons not from blackboxes will be numerically inverted for solution.
- Integrate** Invoke the selected integrator (LSODE currently available) on the problem.

### 7.1.4 SOLVER ANALYZE MENU

- Reanalyze** Reanalyzes the current problem.
- Debugger** Opens a tool which deals with the system as a numbered list of variables and relations. See Section 7.4, "Debugger," on page 64 for more information about the Debugger.

**Overspecified** Finds and displays the variables that can be freed to reduce the degrees of freedom in an overspecified system.

**Find dependent eqns.** Finds structural or numeric dependencies of a system.

**Numeric Dependency.** Doesn't mean much on an unsolved system. This command inverts one block at a time and checks the blocks for numeric dependency using the QRS1v solver. Any non-zero dependency is reported, but those relations with coefficients down around machine epsilon (1e-16) are probably not dependent. Poorly scaled problems can appear more singular than they really are.

**Structural Dependency.** Find the equations or variables involved in a structural dependency. For systems that should be square, this is similar to overspecified, but for DAE's this detects the equations which need to be differentiated according to Pantelides. The user interface for reporting the data returned is not complete.

**Find unassigned eqns.** Shows the equations which cannot be assigned by the structural analysis.

**Evaluate unincluded eqns.** Evaluates the residuals of unincluded relations and checks them for convergence. This may not be a wise idea, depending on why the relations have been excluded.

**Find vars near bounds** This will write variable names passing test

$$\text{abs}(\text{value-bound})/\text{nominal} < \text{epsilon} \quad (7.1)$$

to the console. The test is performed first for lower bounds and then for upper bounds and the results are clearly marked. This can be used for locating variables which may yield a more tractable problem when moved to the bound and fixed while freeing another variable. The value of Epsilon can be set from the Solver's General parameter page.

**Find vars far from nom** This will write variable names passing test

$$\text{abs}(\text{value-nominal})/\text{nominal} > \text{bignum} \quad (7.2)$$

to the console. This test can be used for locating variables which are poorly scaled and for evaluating where model initialization methods need improvement. The value of bignum can be set from the General parameter page.

### 7.1.5 SOLVER EXPORT MENU

- to Browser**            This button sends the instance currently in the Solver to the Browser.
- to Probe**              This button sends the instance currently in the Solver to the Probe piecewise, that is all the variables and relations get shipped, not the instance name itself.

## 7.2 SOLVER BUTTON BAR

The solver button bar, which is located just below the solver menu, contains three buttons, the solver select button, the solver options button, and the halt button.

- Solver Select Button**            This button contains the name of the currently selected solver. Depressing this button reveals a menu of available solvers which can be selected.

- Solver Options Button**            The Options menu on the solver allows the user to view and to change the settings for the parameters associated with ASCEND's solvers. A solver's parameters may be changed even when the solver is empty of another solver is selected. Depressing the options button reveals a list of parameter pages which can be selected for viewing (and editing).

Below, we discuss using the parameter pages and the general solver parameters; solver specific parameters are discussed below in Section 7.3, "Available Solvers," on page 62.

- Halt Button**                    Halts the solver and returns control to the interface as soon as possible. Not all solvers connected to ASCEND will respond to the halt signal.

### 7.2.1 GENERAL PARAMETERS PAGE

Selecting General under Options will display the General Parameter Page (See Figure 7-3). This is where we keep items relevant to the interface and to the way mathematical specialty functions and utilities are handled in ASCEND. Following, we will discuss the parameters that appear on this page.

Figure 7-3 General Parameter Page

iterations before screen update	10
cpu sec before screen update	3
modified log epsilon	1e-08
bound check epsilon	1e-3
far from nom bignum	10e3
integrator state log	y.dat
integrator observation log	obs.dat
integrator log SI units	display
integrator log columns	variable
<input checked="" type="checkbox"/> overwrite integrator logs	
<input type="checkbox"/> check numeric rank after solving	
<input type="checkbox"/> show block summary	
<input type="button" value="OK"/> <input type="button" value="Help"/>	

Iterations before screen update

Because the interface update is sometimes rather time consuming (or more accurately when the window manager is slow, the interface holds up the solver) this specifies how many iterations to stay down in the solver algorithm before returning to the user interface to update statistics. In the case of floating point errors or solution completion before the limit is reached, the return and update will happen immediately rather than waiting for the limit to be reached. For solvers that don't truly iterate in an accessible fashion (e.g. MINOS) this parameter is ignored.

CPU seconds before screen update

For solvers which do offer access to status information between iterations, this is the maximum number of cpu seconds before an interface update. If, while still not done with the number of iterations given in "iterations before screen update," the solver algorithm detects that the cpu seconds limit has expired, then it will return early to update the interface. At least one iteration will be completed before the clock is checked.

Modified log epsilon

This parameter controls the value for epsilon in the "lnm" function. Lnm can be used instead of natural log (ln) when the argument is likely to be very small or to go negative in the solution process. This avoids a host of floating point errors in initialization and solving of many kinds of models.

The modified natural log function  $f$  is defined as

$$f(x) = \begin{cases} \ln(x) \forall (x > \epsilon) \\ \frac{x}{\epsilon} - 1 + \ln(\epsilon) \forall (x \leq \epsilon) \end{cases}$$

The first derivative of this function is continuous. The second derivative has a jump from 0 to  $-1/\epsilon^2$  at  $x = \epsilon$ .

<u>Bound check epsilon</u>	This is the epsilon parameter used in the [link: to obvious location/Find vars near bounds] utility under the Solvers Analyze Menu.
<u>Far from nom bignum</u>	This is the bignum parameter used in the [link: to obvious location/Find vars far from nom] utility under the Solvers Analyze Menu.
<u>Integrator state log</u>	This is the name of the file for integrator state variable output. It defaults to y.dat in the current directory.
<u>Integrator observation log</u>	This is the name of the file for user defined observation output during integration. It defaults to obs.dat in the current directory.
<u>Integrator log SI units</u>	This switch causes the output to be written in SI units or in the user's selected interface units.
<u>Integrator log columns</u>	This option selects how the state and observation logs should be formatted. We can produce fixed or variable width formats suitable for import into nearly any other software package.
<u>Overwrite integer logs</u>	This switch lets the user control whether integration log files should be appended or replaced with each run.
<u>Check numeric rank after solving</u>	When selected the numeric rank will be checked at the solution and a message will be displayed if the system is rank deficient.
<u>Show block summary</u>	When selected the cost statistics (cpu, iterations, evaluations) for all blocks of significant size will be listed to the screen after each solve.

## 7.3 AVAILABLE SOLVERS

Here is the list of solvers that at one time or another have been connected to ASCEND:

- Slv

- QRSlv
- LSODE
- MINOS
- LSSlv
- Opt (SQP)
- CONOPT
- Make MPS

All of these solvers may not be available in your installation of ASCEND. A brief description of ASCEND's primary solver, QRSlv, follows.

### 7.3.1 QRSLV

QRSlv is a nonlinear algebraic equation solver based on the paper "A Modified Least Squares Algorithm for Solving Sparse NxN Sets of Nonlinear Equations" by A. Westerberg and S. Director (EDRC TECH REPORT 06-5-79).

#### 7.3.1.1 PARAMETERS

Following is an incomplete list of control parameters for the QRSlv algorithm. Most users will only change the time limit, iteration limit, and maximum residual as the default parameter values work quite well.

<u>Time limit</u>	The total number of seconds allowed in 1 push of the Solve button.
<u>Iteration limit</u>	Total number of iterations in for any single partition in the problem.
<u>Minimum pivot (epsilon)</u>	the smallest pivot value allowed in the linear solution of a subproblem.
<u>Pivot tolerance</u>	pivot selection criterion.
<u>Maximum residual</u>	This is the maximum absolute error that QRSlv is allowed to consider an equation as solved. Self scaling equations will more easily satisfy this than those that aren't. E.g. an energy balance (with terms the size of $10^8$ ) will have a far harder time meeting this convergence criterion if you do not divide them through by an appropriate constant.
<u>Partitioning</u>	If off, entire problem will be solved as a block. Divergence is usually the result on nonlinear problems of any size above 25 or so.

<u>Detailed info</u>	QRSlv spews all sorts of info if you turn this switch on. The utility of such info is often as much for the authors of slv as for the user. The volume of info is large. Most of the spew (that to do with singletons (1x1 blocks) is suppressed if the switch ‘show singletons details’ is off.
<u>Auto-resolve</u>	When complete, the solver is supposed to rerun itself for changes of significance made in the interface if this switch is on.
<u>write to file</u> <u>SlvLinsol.dat</u>	If this switch is on, a whole set of files named SlvLinsol.dat.X where X is integer are produced during the solution of the problem. The X increments for each successive linear system inversion or solution. The files contain Jacobian and rhs data in machine readable forms for import to stand-alone solver tools. There are generally quite a lot of them. X always starts at 0 for a given ascend session and goes up from there.
<u>show singletons</u> <u>details</u>	When the ‘detailed solving info required’ switch is ON this switch controls whether or not full singleton solving information is shown. In particular, if this is off all direct solve spew is cancelled, leaving that which usually of interest, the NxN block solution iterations, to be displayed.
<u>bipartial pivoting</u>	An experimental option for stabilizing the RANKI algorithm on hard problems. It enables searching of both current row and column during linear factorization. It is somewhat more expensive in terms on fill and CPU time, but can lead to solution of otherwise unsolvable problems. The modification is due to Joe Zaher. This option is likely to be replaced by a choice of several linear routines eventually. The original motivation came from distillation models which become illconditioned as tray number grows.

## 7.4 DEBUGGER

The Debugger shown in Figure 7-4 is an aid for examining the variables and relations in the Solver. The debugger is often used in tandem with the incidence matrix because the debugger is queried using the solvers’s internal relation/variable indexing (which starts at 0). When a variable (relation) number is typed in the variable (relation) entry box the variable (relation) Name and Attribute buttons may be clicked to obtain information about the variable (relation). The information is printed to the console window. The variable (relation) may also be exported to either the Browser or Probe by making the appropriate selection under the export pull down menu.



Figure 7-4 The Debugger Window

Variable:
<input type="text"/>
Name
Attributes
Export
Equation:
<input type="text"/>
Name
Attributes
Export
Block:
<input type="text"/>
Variables
Equations
Export to probe
System:
Variables
Export to probe
<input type="button" value="OK"/>
<input type="button" value="Help"/>

When a variable or relation number is entered in the debugger, the corresponding partitioned block number appears in the 'block' entry box. Statistics on the number of rows and columns in the block are displayed just below the block entry box. Note that a block number can also be entered directly into the block entry box. The Variables (Equations) pull down menu below the block entry box contains the selections Values, Attributes, and Probe (and Find Dependent). Selecting Values or Attributes will write the requested information to the console for each variable (equation) in the block. Selecting Probe will export the block's variables (equations) to the probe. Selecting Find Dependent under the Equations pull down menu will write the name of any dependent equations within the block to the console. Selecting the Export to Probe button will export both the block's variables and equations to the probe.

The debugger also responds to requests relating to the overall system, or current solver instance. A Variables pull down menu and an Export to Probe button are located beneath the 'System' label on the debugger. The Variables menu contains the selections Values, Attributes, Reset Values, and Reset Nominals. Selecting Values or Attributes will write the requested information to the console for each variable in the system. Selecting Reset Values will reset the system's variables to their nominal values. Selecting Reset Values will reset the system's nominals to their current variable values.

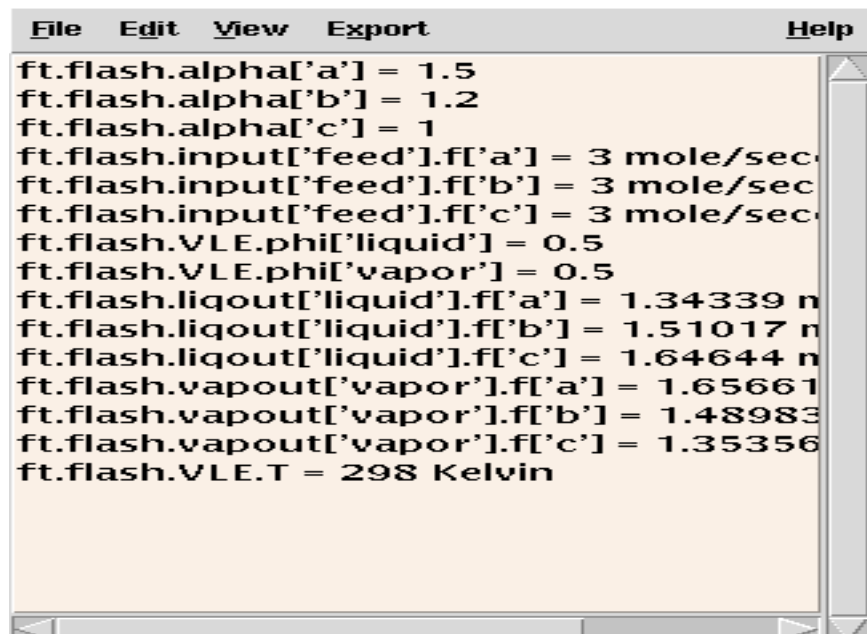


# CHAPTER 8 THE DATA PROBE WINDOW

## 8.1 OVERVIEW

The [data probe shown in Figure 8-1](#) is a window which manages collections of names from the ASCEND simulation universe. Each collection is kept in a buffer, and the user can switch among as many buffers as are needed for convenience. For example, the first buffer may be used as a set of bookmarks to store the names of interesting submodels within a large simulation, a second buffer can be used to monitor a set of key variables, and a third can be used to monitor specifications. The browser provides a two-level view of information - the probe provides a random access view.

Figure 8-1 Probe window



```
File Edit View Export Help
ft.flash.alpha['a'] = 1.5
ft.flash.alpha['b'] = 1.2
ft.flash.alpha['c'] = 1
ft.flash.input['feed'].f['a'] = 3 mole/sec
ft.flash.input['feed'].f['b'] = 3 mole/sec
ft.flash.input['feed'].f['c'] = 3 mole/sec
ft.flash.VLE.phi['liquid'] = 0.5
ft.flash.VLE.phi['vapor'] = 0.5
ft.flash.liqout['liquid'].f['a'] = 1.34339 n
ft.flash.liqout['liquid'].f['b'] = 1.51017 n
ft.flash.liqout['liquid'].f['c'] = 1.64644 n
ft.flash.vapout['vapor'].f['a'] = 1.65661
ft.flash.vapout['vapor'].f['b'] = 1.48983
ft.flash.vapout['vapor'].f['c'] = 1.35356
ft.flash.VLE.T = 298 Kelvin
```

Names are imported to any collection buffer from the other parts of the user interface or from a previously saved file of names. Once collected, a name remains in the buffer until the user removes it, even if the type library and simulations are deleted. This way the set of names is preserved when the user makes a small modification to a MODEL and rebuilds it.

Names in probe buffers are displayed with their corresponding values or other attributes as appropriate. When a name is not well defined (perhaps because the simulation it came from has been deleted temporarily) the attribute displayed is "UNCERTAIN." As soon as the name becomes well-defined again by having a corresponding simulation object built, the correct attribute will appear. Names of atomic objects (reals, integers, sets, symbols, booleans) which have not yet been assigned a value will be shown as "UNDEFINED" until some operation assigns them a value.

## **8.2 THE FILE MENU**

### **8.2.1 NEW BUFFER**

This starts another collection of names, which is initially empty. Each buffer receives a standard name when it is created, NoNameX.a4p, where X is the number of the buffer. These buffer names appear at the bottom of the File menu.

### **8.2.2 READ FILE**

This appends a file full of names into the current buffer and will automatically attempt to associate them with the simulations in the system. This way the name list can be reloaded from a prior work session. The file name is not associated with the buffer.

### **8.2.3 SAVE**

This will save the names in the current buffer to a file with the buffer's menu name. If you wish to save with a more meaningful name, use "Save as." Values are not saved with these names. To save the values, use the Print command.

### **8.2.4 SAVE AS**

This allows you to specify the directory and file name in which to save the names in the current buffer.

### **8.2.5 PRINT**

This lets you print the current buffer to a printer or a file. This prints what you see in the buffer window, including the values. The printer setup dialog will pop up for you to set the destination.

## **8.3 THE EDIT MENU**

### **8.3.1 REMOVE SELECTED NAMES**

This options removes all highlighted lines in the window. The selection in the probe can be set in a disjointed fashion using Control-Button-1 and drag.

### **8.3.2 REMOVE ALL NAMES**

This options removes all names in the current buffer window.

### **8.3.3 REMOVE UNCERTAIN NAMES**

This removes all names that are not well defined. These are the names displayed as “name = UNCERTAIN.”

### **8.3.4 COPY**

This copies all the selected items in the current buffer to the clipboard.

## **8.4 THE VIEW MENU**

From the view menu one can select a new font for the probe by the standard font dialog, and one can toggle the Probe’s auto-display feature.

## 8.5 THE EXPORT MENU

### 8.5.1 TO BROWSER

This option sends the first selected name in the probe to the browser.

### 8.5.2 TO DISPLAY

This options sends some form of the selected names in the probe to the Display slave window, replacing whatever used to be in the display.

## 8.6 THE PROBE FILTER

A class or classes of object can be imported to the probe en masse. The [import filter shown in Figure 8-2](#) lets you select which collection of names (probe buffer) is to receive the imported names which are of the types checked. Currently the probe filter window is accessible only from the Browser Export button.

The filtering import can also be executed from the Script using the PROBE command. The list of ones and zeros required for the PROBE command is ordered in the same way as the list of types in the import filter window. The easiest way to set the list of ones and zeros is to use the Script recording feature and the Browser Export Many to probe button.

Figure 8-2 Probe import filter

Exporting from: tvlf.Feed

Probe buffer: current

- Collect relations
- Collect logical relations
- Collect booleans
- Collect integers
- Collect reals
- Collect symbols
- Collect sets
- Collect subatomic booleans
- Collect subatomic integers
- Collect subatomic reals
- Collect subatomic symbols
- Collect subatomic sets
- Collect boolean constants
- Collect integer constants
- Collect real constants
- Collect symbol constants

OK Help Cancel



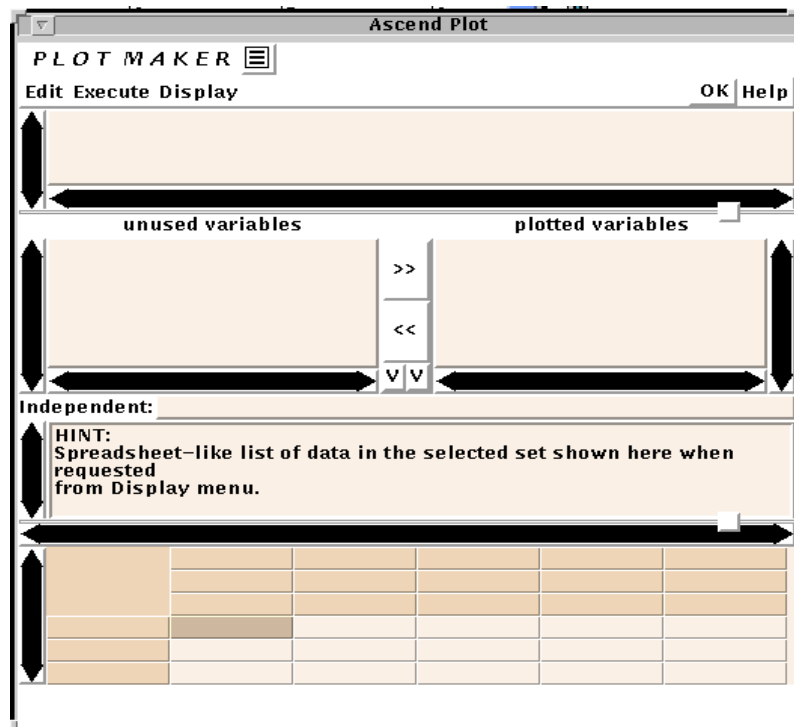


# CHAPTER 9 ASCPLOT

## 9.1 PLOT MAKER

The following contains a description of the options available in each of the *Ascend Plot* menus. The *Ascend Plot* window shown in Figure 9-1

Figure 9-1 The Ascend Plot Window



is a result of clicking on the ascplot button from the Toolbox with the left mouse button.

### 9.1.1 THE EDIT MENU

From the **Edit** Menu, the following options are available when a data set has not yet been loaded: **Load data set** and **Select grapher**. The **Save data set**, **Unload data set**, and **Merge data sets** options are available after one or more data sets have been loaded into the plot window.

#### 9.1.1.1 LOAD DATA SET

Selecting **Load data set** opens the *File select box* window. This window is used to select the file that contains the data generated from the dynamic simulation. The default file is obs.dat. This file contains the observation variables as set forth in the dynamic library models. After having selected the appropriate file, press the OK button and return to the *Ascend Plot* window.

#### 9.1.1.2 SAVE DATA SET

This option is currently not functional.

#### 9.1.1.3 UNLOAD DATA SET

By highlighting the desired data set and selecting **Unload data set** from the **File** menu, the user can remove the data set from the *Ascend Plot* window. The *Delete these data sets?* window appears to verify that the user wants to remove the indicated data sets.

#### 9.1.1.4 MERGE DATA SETS

#### 9.1.1.5 SELECT GRAPHER

Currently, the only supported grapher is Xgraph (or its tk flavored version tkxgraph). Other possible graphers are XMGR and gnuplot. Since these graphers are not distributed with the ASCEND distribution, they are also not supported.

### 9.1.2 THE EXECUTE MENU

To plot the variables in the plotted variables section, select **View plot file** from the **Execute** menu.

### 9.1.2.1 VIEW PLOT FILE

This option will plot the variables displayed in the plotted variables section of the *Ascend Plot* window.

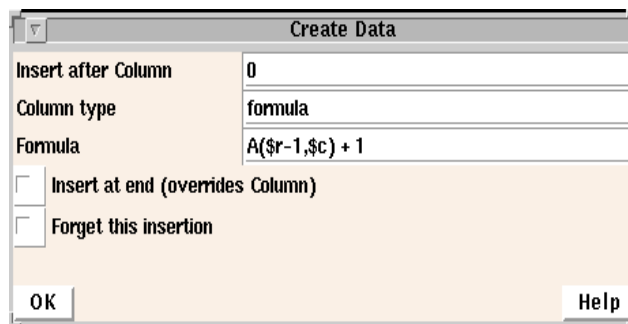
### 9.1.2.2 WRITE PLOT FILE

To save the output in its graphical representation, select **Write plot file** from the **Execute** menu. Selecting this option opens the *File select box*. Enter the name of the file to be saved and press the OK button. The default extension for the graph is .xgraph.

### 9.1.2.3 INSERT COLUMN

Selecting the **Insert column** option from the **Execute** menu opens the *Create Data* window. This window is shown in Figure 9-2.

Figure 9-2 The Create Data Window



There are several options available from the *Create Data* window.

#### 9.1.2.3.1 Insert after Column

This can be any number between 0 and the maximum number of variables in the observation file. For example, if the user wishes to add a column after the third column, the user should enter a 3 in this space.

#### 9.1.2.3.2 Column type

The default value for Column type is data, however by placing the cursor over the data box and pressing the left mouse button, another option is revealed. The other option is formula. The user should select data if no formula can be used to describe the information to be added to the spreadsheet. The user should select formula if that is appropriate. In this case, a column was inserted after Column 0 and we are using the formula Column type.

### 9.1.2.3.3 Formula

If the data option was selected in the previous section, then this does not apply. However, if the formula option was selected, then the user can edit the default formula. The default formula takes the value of the variable in the current row ( $r$ ) and the column before the new column ( $c-1$ ) and adds one ( $+1$ ) to it.

### 9.1.2.3.4 Insert at end (overrides Column)

The user can select this box to place the new column after the last column in the spreadsheet. This will override anything in the Insert after Column line.

### 9.1.2.3.5 Forget this insertion

The user can select this box to ignore the changes made to the spreadsheet.

### 9.1.2.4 RECALCULATE COLUMN

This option is currently not functional.

### 9.1.2.5 INSERT ROW

The insert row option has the same options as the Insert Column option. Note that the formula take the value from the row immediately before it ( $r-1$ ) and the current column ( $c$ ) and adds one ( $+1$ ) to it.

### 9.1.2.6 RECALCULATE ROW

This option is currently not functional.

## 9.1.3 THE DISPLAY MENU

The **Display** menu has various features which include showing and hiding the data in the spreadsheet, setting plot titles, loading old plots, updating existing plots, and deleting plots.

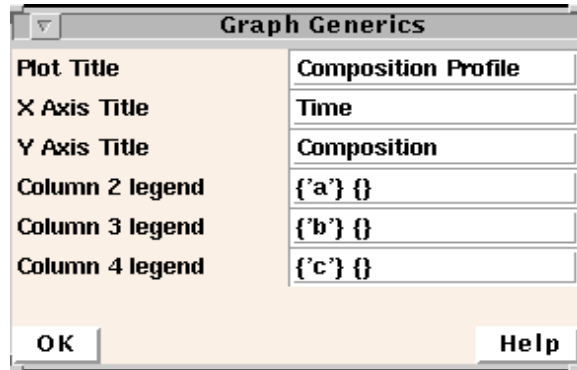
### 9.1.3.1 SHOW DATA / HIDE DATA

Selecting the **Show data** option from the **Display** menu loads the data into the spreadsheet in the bottom section of the *Ascend Plot* window. This option then toggles to **Hide data**. Selecting this option will hide the data just loaded into the spreadsheet section of the window.

### 9.1.3.2 SET PLOT TITLES

Selecting the **Set plot titles** option from the **Display** menu opens the *Graph Generics* window. This window is shown in Figure 9-3.

Figure 9-3 The Graph Generics Window



There are several options within this window depending on the number of variables being plotted.

#### 9.1.3.2.1 Plot Title

The user can change the default title (AscPlot) to something that is more descriptive and meaningful for the given data. In this case, we set the title to be Composition Profile since we are plotting the mole fractions of the components in the system.

#### 9.1.3.2.2 X Axis Title

The user can change the default title (X) to something more descriptive. In this case, we are plotting the time on the x-axis.

#### 9.1.3.2.3 Y Axis Title

The user can change the default title (Y) to something more descriptive. In this case, we are plotting the Composition on the y-axis.

#### 9.1.3.2.4 Column # legend

In this case, (#) is the number of the variable being plotted. If variables 2, 3, and 4 are being plotted, they will be entries in the *Graph Generics* window entitled Column 2 legend, Column 4 legend, and Column 4 legend. These entries can be changed to something less descriptive than the default. Usually the default for this field is a bit much. In this case, the legend was changed to 'a', 'b', and 'c'.

### 9.1.3.3 LOAD OLD PLOT

This option is currently not functional.

### 9.1.3.4 UPDATE PLOT

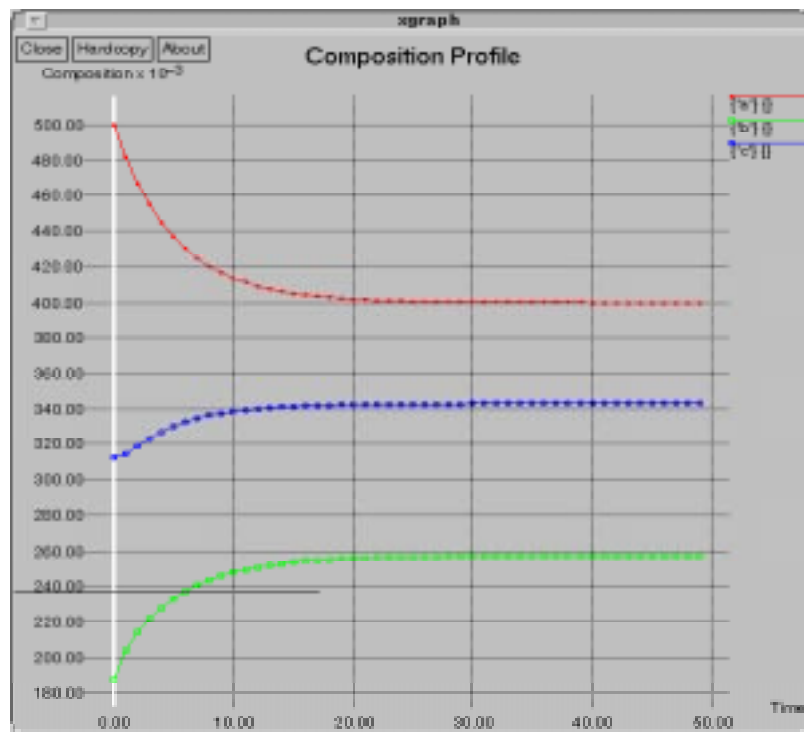
This option is currently not functional.

### 9.1.3.5 DELETE PLOT

This option is currently not functional.

The plot of the completed graph is shown in Figure 9-4.

Figure 9-4 Complete Plot



### 9.1.3.6 THE GRILL



The grill is located directly to the right of PLOT MAKER. Clicking on this button with the left mouse button opens the *XGraph Control* window. By clicking on the More button located at the bottom of the window, the user can scroll through numerous available options for the graphs. Some of these options include line color, fonts, graph type (i.e. log or semilog), and marker types. These are left to the user to explore.

## 9.2 NAVIGATION

Open a file using the **Load data sets** option from the **File** menu. You will notice that the selected file is now displayed in the top section of the *Ascend Plot* window. By double-clicking on the file name with the left mouse button, the observation variables are now placed in the section entitled unused variables. The unused variables are the list of variables that the user does not want to look at in the current graph.

To select a variable to plot, highlight the desired variables using the left mouse button and click on the (>>) button. This will move the variable from the unused variables list to the plotted variables list. Once this is done, you can now plot the variable.

The two buttons separating the unused variables section and the plotted variables section are used to add (>>) and remove (<<) variables to and from the plotted variables list.

You will notice that there is a section of the *Ascend Plot* window entitled Independent. Here the independent variable is time. This was set in the dynamic library file. If the user desires to look at the phase plot of two of the compositions, the user must move one of the compositions into the Independent variable position.

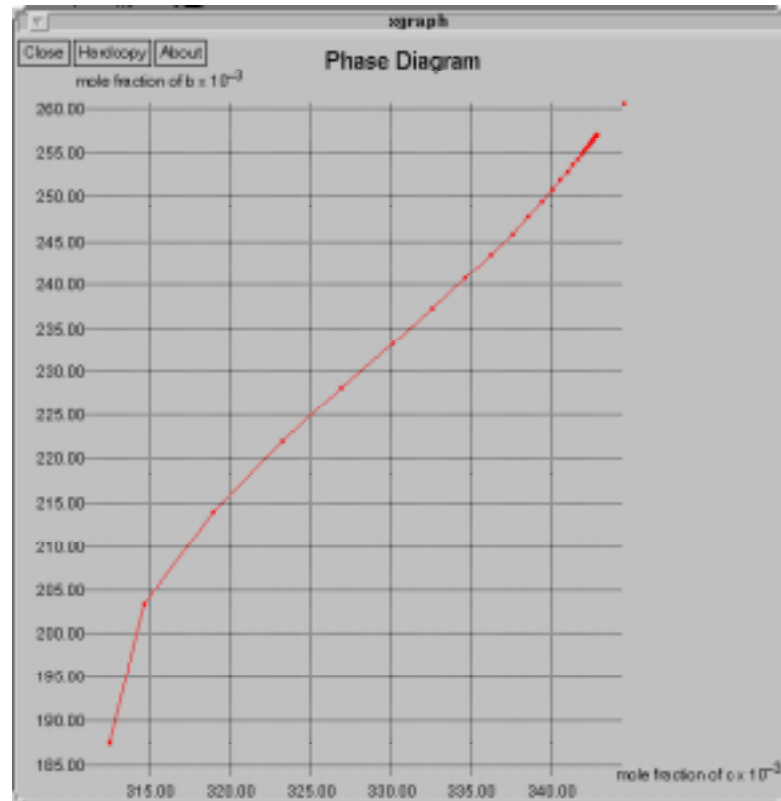
To do this, let's assume that all of the variables are currently in the unused variables list and we wish to plot the composition of component 'c' versus the composition of component 'b'. Thus, component 'c' is now going to be our independent variable. Highlight component 'c' in the unused variables list and press the (V) button. This button is one of two buttons located directly under the (>>) and (<<) buttons. The (V) button on the left is used to move variables between the unused variables list and the Independent variable list while the (V) button on the right is used to move variables between the plotted variables list and the Independent variable list. Therefore, we are going to use the (V) button on the left.

By doing this, we see that the composition of component 'c' is now the independent variable and the time is now an unused variable. Select the composition of component 'b' and press the (>>) button to move the variable from the unused variables list to the plotted variables list. The only remaining task is to edit the plot title and axes using the **Set plot titles** option from the **Display** menu. Assuming we have done this as described above, the resulting graph is shown in Figure 9-5.



The remaining section of the *Ascend Plot* window is the HINT: section. This section contains a brief description of the various buttons and sections of the Ascend Plot window.

Figure 9-5 Phase Diagram

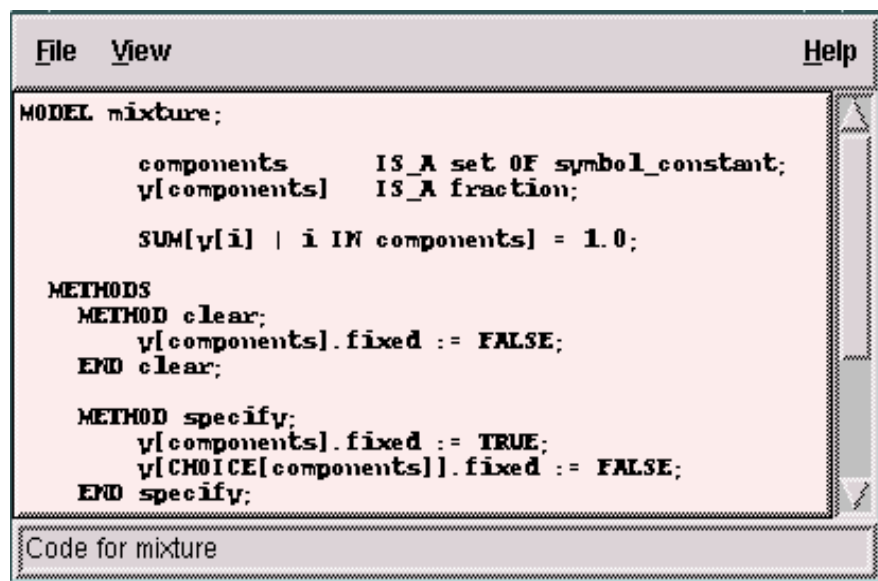


# CHAPTER 10 DISPLAY SLAVE

## 10.1 OVERVIEW

The display slave window (Figure 10-1) functions as a dumping ground for information which is too complex to display in other ways in the Library, Browser or other primary windows. It has rudimentary editing abilities so the user can manually adjust the format of displayed information if needed, for example by rearranging a highly nonlinear relation with more than a few variables. Changes to displayed text do not affect the rest of the system in any way.

Figure 10-1 Display slave window



The screenshot shows a window titled "Display Slave" with a menu bar containing "File", "View", and "Help". The main area contains the following code:

```
MODEL mixture;  
  
    components      IS_A set OF symbol_constant;  
    y[components]  IS_A fraction;  
  
    SUM[y[i] | i IN components] = 1.0;  
  
METHODS  
METHOD clear;  
    y[components].fixed := FALSE;  
END clear;  
  
METHOD specify;  
    y[components].fixed := TRUE;  
    y[CHOICE[components]].fixed := FALSE;  
END specify;
```

At the bottom of the window, there is a text box containing the text "Code for mixture".

## 10.2 THE FILE MENU

### 10.2.1 PRINT

This option brings up the default print dialog described in the section Utilities. The print command can be used to save the displayed text to a file.

### 10.2.2 CLOSE WINDOW

The option closes the display window.

## 10.3 THE VIEW MENU

### 10.3.1 SHOW COMMENTS IN CODE

This option controls whether or not comments are displayed when code is displayed **as read from source files**. This setting is not retroactive; that is code already displayed will not be redisplayed when changing this setting.

When code is displayed in the machine representation, i.e. with equations and set expressions in postfix (reverse polish) notation, comments are never displayed.

### 10.3.2 FONT

This option brings up the standard font setting dialog so you can change the size, style, and font of the characters in the display window.

### 10.3.3 OPEN AUTOMATICALLY

This option controls whether or not the display slave window opens automatically when it receives information. Sometimes it is easier to send several items to the display and then open it at the end.

## 10.4 TITLE LINE

The title line at the bottom of the window is set by the last client to export something to the display. The user may edit the title, but the next time new information is displayed, these edits will be lost.

# CHAPTER 11 ASCEND UNITS

## 11.1 THE MENU BAR

The Units Tool Set provides tools to allow the user to change the display units for variables.

### **Units vs dimensions**

We distinguish between *units* and *dimensions* in ASCEND. The dimensions of acceleration, for example, are  $L/T^2$ , i.e., length/time squared. Units for acceleration are:  $m/s^2$ ,  $ft/hr^2$  and so forth.

### **Typical use**

The user will typically first pick the overall system of units such as SI, American Engineering or cgs. Alternatively the user may select to use the *default* display of units for some or all variable types. Displaying in default units means ASCEND will present the units in terms of the ten basic dimensions supported by ASCEND (length, time, temperature, etc.). The user can select the units to be used for each basic dimension. Whichever of these alternatives the user selects, he or she may then also choose the units ASCEND should use to display particular variable types. An example would be to select first SI units, then override the display of energy to be in default units and pressure to be in atm.

Once users have created their favorite choices for display units, they may save them to files for later restoration.

We describe here the various tools available within the Units tool set.

### 11.1.1 UNITS EDIT MENU

- Set precision** Use the slider switch for this tool to set the number of digits of precision for displaying variable values to between 4 and 16. Precision is the number of digit displayed when the number is displayed using scientific notation. For example, 0.12345678 e04 for 1234.5678 has a precision of 8 digits.
- Read file** Reads in a file previously saved using the “Write file” command. Restores the display units to those previously saved.
- Write file** Writes out (in the current working directory) a plain text version of the user specified display units. Units which are defaulted are not written to this file. One can restore the display units to those currently set by reading this file back in later.

### 11.1.2 UNITS DISPLAY MENU

- Show all units** Causes the Display window to open showing all the units conversions currently used in ASCEND.
- SI (MKS)** Pushing this button makes the default display units SI units.
- US Engineering** Pushing this button makes the default display units US Engineering units.
- CGS** Pushing this button makes the default display units CGS units.

### 11.1.3 UNITS HELP MENU

- An essay on units vs dimensions** ASCEND stores all numbers in SI (MKS) units internally. The units associated with a dimensionality (as exemplified by some atom) will be used when displaying variables of that dimensionality. These units can be manipulated through the Units window.

Numbers with unrecognized dimensionality (higher derivatives, multipliers, residuals and what not) will be given units consistent with the display units defined for the 10 base dimensions. The display units for the 10 dimensions can be changed through the Units window Display menu if you prefer an alternate default set such as US engineering, and so forth.

We recognize 10 base dimensions in the compiler:

L distance meter m

M	mass	kilogram	kg
T	time	second	s
E	e- current	ampere	A
Q	quantity	mole	mole
TMP	temperature	Kelvin	K
LUM	luminous intensity	candela	cd
P	plane angle	radian	rad
S	solid angle	steradian	srad
C	currency	currency	CR

The units conversions are defined in \$ASCENDDIST/compiler/units\_input, which is not particularly restricted. Units\_input is converted to an efficient binary form (unitsfile.uni) at the time ASCEND is installed.

It can be argued that C is not a fundamental dimension, from a physical standpoint. There is more to life than physics: there is economy, hence engineering, hence an Advanced System for Computations in ENgineering Design.

The dimensions P and S are ‘supplementary’ according to the General Conference, but their use makes the coding of ASCEND much cleaner and easier.

## On UNITS

The left box in the Units window lists a set of atom types, each having a unique dimensionality. Selecting an atom in the left box will fill the right box with different possible units that the system knows about to display this type of variable. Dimensionless atoms and wild dimensioned atoms are not shown since they do not have display units. If you do not see an atom you expect here, it is because ASCEND already found another atom of the same dimensionality, e.g. fugacity may show up instead of pressure.

Selecting a unit in the right box sets that unit as the display unit for all variables having the same dimensionality of the selected atom in the left box. Thus picking **atm** for fugacity will also change pressure units to **atm**. Selecting ‘default’ will cause the display to be a combination of

the *fundamental* units (a nice way to remind oneself of the fundamental units for energy, for example).

Fundamental units are the units corresponding to single dimensions. These units are chosen on the Display menu under the dimension choices. No atoms with fundamental units are listed in the left box. The current set of fundamental units is always shown at the very bottom of the units window. This set is used whenever a value is displayed which does not have a user specified units set associated with its dimensionality. The fundamental units are created via the `units_input` file mentioned above. If you do not find one you want, ask whoever compiled your version of `unitsfile.uni` to add the missing unit and rebuild the `unitsfile.uni`.

If converting the units for a variable makes the display of that number impossible (e.g., due to overflow). ASCEND will first attempt to display it using its fundamental units. If it still cannot be displayed, it will be displayed in SI units.

You may specify a new combination of existing units (e.g. Pa\*s) using the **Set units** which is the line at the bottom of the window. Type in the combination desired and press RETURN.

Unit strings may not have parentheses in them. For example, kg/(m\*s<sup>2</sup>) is not allowed.

# CHAPTER 12 THE ASCEND TOOLBOX

The [toolbox window shown in Figure 12-1](#) lets the users open and close the various windows for the tool sets available in ASCEND. The toolbox window is a vertical window containing 12 buttons: exit, ascplot, help, utilities, bug report, LIBRARY, BROWSER, SOLVER, PROBE, UNITS, DISPLAY, SCRIPT.

The buttons in the toolbox with names in ALL CAPS (LIBRARY and following) open and close the windows for the corresponding tool sets

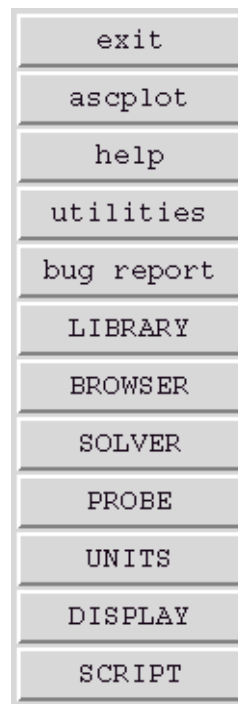


Figure 12-1 The ASCEND Toolbox window.



in ASCEND<sup>1</sup>. As each of these toolsets has its own documentation, we shall not discuss them here. We discuss here only the first set of buttons: Exit, Ascplot, Help, Utilities, and Bug Report.

## 12.1 EXIT

This button shuts down all of ASCEND after making sure you want to quit. ASCEND does no checking to see if there is unsaved work so be certain you have saved what you want of it before selecting this button.

(For more advanced users, we note that, just before exiting, we call the tcl function `user_shutdown` which may be redefined in the `.ascendrc` file in your HOME directory. Under Windows, the `_ascendrc` is the name of the corresponding file.)

## 12.2 ASCPLOT

Selecting this button opens the plotting tool for ASCEND. You can find any file that contains data for a plot and plot it with this tool. Ascplot is described elsewhere.

## 12.3 HELP

Pressing this button will provide access to the Help Documentation for ASCEND. The help system is described elsewhere.

## 12.4 UTILITIES

Selecting this button opens the system utilities window. The system utilities window is described elsewhere.

## 12.5 BUG REPORT

The link  
<http://www.cs.cmu.edu/~ascend/Email.html>  
is connected to the web server for ASCEND at CMU. Alternatively, send a bug report to

---

1. The more advanced user should note that changing the `iconname` of a window (via `ascend.ad`) does not change its toolbox name.

`ascend+bugs@cs.cmu.edu`

if you cannot access this link. We do not have an 800 number, but we usually get to bug reports very quickly.

When submitting a bug report, please try to

1. Duplicate the error.
2. Tell us in excruciating detail how you duplicated it.
3. Report to us the platform and operating system (OS) on which you are running. Also please tell us the distribution number for the ASCEND code on which you are running. This is research software. We are not committed to backward compatibility, and we do not have access to all the platform/OS combinations out there. If the bug you report has been fixed in a newer version, your only fix is to get the new version or fix it yourself. If you are on a platform to which we do not have access, we will consider working out the bugs with you in the hope that you will then give us back a copy for the new platform.
4. Send along any model code you have that is involved in the bug manifestation. It may happen that, in the process of fixing the ASCEND bug, we could fix some of your model bugs. We are not in the business of debugging your model code unless it is also interesting to our research. We often find new applications of ASCEND interesting, however.
5. Subscribe to the ASCEND user mailinglist/bboard: Send mail to `ascend+subscribe@edrc.cmu.edu`.



# CHAPTER 13 THE SYSTEM UTILITIES WINDOW

## 13.1 OVERVIEW

The [system utilities window shown in Figure 13-1](#) displays and allows modification of the variables which control the interaction of ASCEND with the operating system and with other programs.

The values of the variables are initialized from the user's environment, from the file `.ascend-config` in the user's HOME directory, and from settings within ASCEND.

If the user chooses to save the system utility settings, ASCEND writes the current values of the variables into file `.ascend-config` in the user's HOME directory. ASCEND will automatically reread those values in the next time it starts.

ASCEND Utility Settings				
Text print command	>> /dev/null cat			
PRINTER	foo_bar			
TCL_LIBRARY	/usr/local/lib/ascend/lib/			
TK_LIBRARY dir	/usr/local/lib/ascend/lib/			
WWW restart	rMozilla %U			
WWW root URL	/usr/local/lib/ascend/help			
WWW startup	rMozilla %U			
Spreadsheet command	unset			
OK	Save	Read	More	Help

Figure 13-1 The System Utilities window manages ASCEND's interaction with the operating system and with other programs.

When working with the system utilities window, it is important to remember that changes to the variables propagate *immediately* throughout ASCEND, and that there is no way to undo or cancel changes made to the variables<sup>1</sup>.

## 13.2 VARIABLES

The system utilities window contains the following settings. Settings marked with an asterisk\* are not saved in `.ascend-config`.

To change a variable's value, click in the box to the right of the variable's label and type the new value. This new value is immediately available to the ASCEND system.

### 13.2.1 WWW ROOT URL

ASCEND distributes its help system as HTML documents, and spawns a web browser to view these documents. The variable WWW Root URL gives the root of the ASCEND help tree, and the variables WWW Restart Command and WWW Startup Command contains commands to connect to a running web browser and to start a new web browser, respectively.

WWW Root URL contains the first part of a URL to the ASCEND help tree; it is not necessarily a complete URL. The variable should end in a forward slash (/). The **Help** menus and buttons in ASCEND will append text to this value and invoke WWW Restart Command or WWW Startup Command with the complete URL.

The person who installs ASCEND at a site should set this variable to the root of the directory containing that site's copy of the ASCEND help files, for example:

```
file://localhost/usr/local/lib/ascend/help/  
at CMU ICES.
```

The value

```
http://www.cs.cmu.edu/~ascend/help/  
will connect you to the help pages at the ASCEND web site.
```

---

1. Some variables can be restored to the values in effect the last time the system utilities were saved, but this only works if the user has previously saved the values, and it does not restore every variable.

### 13.2.2 WWW RESTART COMMAND

This is a command to redirect the attention of your already running web browser to a new URL. If this command returns an error code, ASCEND will attempt to start a new browser using the WWW Startup Command.

If your favorite browser does not support restarting, set the value of this variable to `false`. This will cause a new browser to start for every help query from the ASCEND interface.

ASCEND will replace every occurrence of `%U` in this command with the URL to be viewed. The default value of WWW Restart Command is

```
netscape -remote openURL(%U)
```

### 13.2.3 WWW STARTUP COMMAND

This is a command to start your favorite web browser. This command is invoked if the value of WWW Restart Command is `false` or if attempting to start a browser using that command returns an error code.

ASCEND will replace every occurrence of `%U` in this command with the URL to be viewed. The default value of WWW Startup Command is

```
netscape %U
```

### 13.2.4 ASCENDLIBRARY PATH\*

The ASCENDLIBRARY variable contains a list of directories that the Library and Script tools search to find files containing ASCEND models and scripts.

The format of the directory list should resemble the PATH environment variable for your platform: a colon (:) separated list of directories (using forward slashes) on UNIX, a semicolon (;) separated list of directories (using backward slashes) on Windows.

The ASCENDLIBRARY variable is initialized from the user's environment or from the ASCEND binary; its value is not saved in the user's `.ascend-config` file.

### 13.2.5 SCRATCH DIRECTORY

The scratch directory is used to write the temporary and plot files that ASCEND creates. The temporary files are automatically deleted before you leave ASCEND, but the plot files are not (since people often want to save plots). You should periodically remove any plot files from the scratch directory, else you may slow build up a large collection of past plot files.

Any existing directory you have write access to can be used as the scratch directory. Under UNIX, `/tmp` is the default value of the scratch directory. Under Windows, the directory given in the environment variable `TEMP`, `TMP`, or `TMPDIR` is used as the default value.

### 13.2.6 WORKING DIRECTORY

Typically, this is the directory you start ASCEND from, but it can be any existing directory you have write access to. Our handling of the working directory is a bit “flaky” at the moment because ASCEND’s command line allows the user to change directories without telling the rest of the interface about it. Intermediate files are sometimes written in the working directory.

### 13.2.7 PLOT PROGRAM TYPE

Currently, the only supported plot types is `xgraph plot` (abbreviated `xgraph`). This setting tells the plot window what type of plot file it should generate.

### 13.2.8 PLOT PROGRAM NAME

This is the name of your plotting program. It should accept the plot type listed in Plot Program Type as input.

The default is `xgraph` on UNIX and `tkxgraph` on Windows. Both `xgraph` and `tkxgraph` are available from the ASCEND web site:  
<http://www.cs.cmu.edu/~ascend/>

### 13.2.9 TEXT EDIT COMMAND

This is a command to spawn your favorite text file editor. (Currently, nothing in ASCEND invokes this command.)

The default is `emacs` on UNIX and `runemacs` on Windows.

### 13.2.10 POSTSCRIPT VIEWER

This is a command to spawn a program for viewing Postscript files. (Currently, nothing in ASCEND invokes this command).

The default is `ghostview` on UNIX and on Windows.

### 13.2.11 SPREADSHEET COMMAND

This is a command to spawn your favorite spreadsheet program. (Currently, nothing in ASCEND invokes this command<sup>2</sup>).

### 13.2.12 TEXT PRINT COMMAND

This entry displays the last command generated by the print dialog box. Changing the value of this entry will have no effect on future printing, since the print dialog manages all aspects of printing.

This value is displayed here as a hold-over from previous versions of ASCEND; developers sometimes use it as a check to make sure the print dialog is doing the right thing.

### 13.2.13 PRINTER VARIABLE\*

This entry displays the last printer the user selected in the print dialog box, or the value of the `PRINTER` or `LPDEST` environment variable if the user has not used the print dialog box during this ASCEND session.

Changing the value of this entry will have no effect on future printing<sup>3</sup>, since the print dialog manages all aspects of printing.

This value is not saved in the user's `.ascend-config` file.

### 13.2.14 ASCENDDIST DIRECTORY\*

The value of the `ASCENDDIST` environment variable is the directory containing the installed ASCEND distribution. If a user can see this

- 
2. Nothing invokes this command because there is no ASCEND code that supports it. Someone needs to write code that will write out the desired variables as columns of numbers suitable for importing into any spreadsheet. If you want to be that someone, let us know and we'll be happy to consult. We have some pseudocode for this already; contact us at `ascend@cs.cmu.edu`.
  3. This is not entirely true. This entry will change the value of the `PRINTER` environment variable (but not the `LPDEST` environment variable). Any command you invoke from ASCEND command prompt that depends on the `PRINTER` environment variable will use the value displayed in this entry.



variable inside the system utilities window, it means its value is correct. Changing the value will most likely cause things to break.

The person who installs ASCEND at a site is typically the only person who needs to be concerned with its value.

The ASCENDDIST variable is initialized from the user's environment or from the ASCEND binary; its value is not saved in the user's `.ascend-config` file.

### **13.2.15 TCL\_LIBRARY ENVIRONMENT VARIABLE\***

The value of the TCL\_LIBRARY environment variable is the directory containing the installed `*.tcl` files required by Tcl. If a user can see this variable inside the system utilities window, it means its value is correct. Changing the value will most likely cause things to break.

The person who installs ASCEND at a site is typically the only person who needs to be concerned with its value.

The TCL\_LIBRARY variable is initialized from the user's environment or from the ASCEND binary; its value is not saved in the user's `.ascend-config` file.

### **13.2.16 TK\_LIBRARY ENVIRONMENT VARIABLE\***

The value of the TK\_LIBRARY environment variable is the directory containing the installed `*.tcl` files required by Tk. If a user can see this variable inside the system utilities window, it means its value is correct. Changing the value will most likely cause things to break.

The person who installs ASCEND at a site is typically the only person who needs to be concerned with its value.

The TK\_LIBRARY variable is initialized from the user's environment or from the ASCEND binary; its value is not saved in the user's `.ascend-config` file.

## **13.3 BUTTONS**

The actions associated with the buttons on the system utilities window are:

### 13.3.1 OK

This button closes the system utilities window. Closing will fail if the scratch directory and working directory are not writable by the user.

### 13.3.2 SAVE

This button writes the current value of most of the variables in the system utilities window to a file called `.ascend-config` in your HOME directory<sup>4</sup>. ASCEND will read this file on startup to get your preferred values.

The variables whose names are in ALL CAPS (i.e., ASCENDLIBRARY, PRINTER, ASCENDDIST, TCL\_LIBRARY, TK\_LIBRARY) are **not** saved to `.ascend-config`. These are environment variables that are set as part of the login process. You may change them interactively, but their interactive values are not saved.

### 13.3.3 READ

The button causes the system utilities window to reread the values stored in `.ascend-config` in your HOME directory. This is useful for editing `.ascend-config` outside of ASCEND while running ASCEND, or for verifying that the changes you saved were properly saved.

### 13.3.4 MORE

The button rotates you through the pages of options.

### 13.3.5 HELP

The button should direct your web browser to this document.

---

4. Under Windows, you can set your HOME directory by setting the HOME environment variable by opening the Control Panel, double clicking the System icon, clicking the Environment tab, and adding the HOME variable to the list of user environment variables.



# CHAPTER 14 FONT SELECTION DIALOG

## 14.1 OVERVIEW

The [font selection dialog](#) (Figure 14-1) is used to select the font for the window from which it is opened. There is no way through the interface to change the font for every ASCEND window.

Currently, the fonts you select are not remembered across invocations of ASCEND. This is a feature we will be adding in a future release.

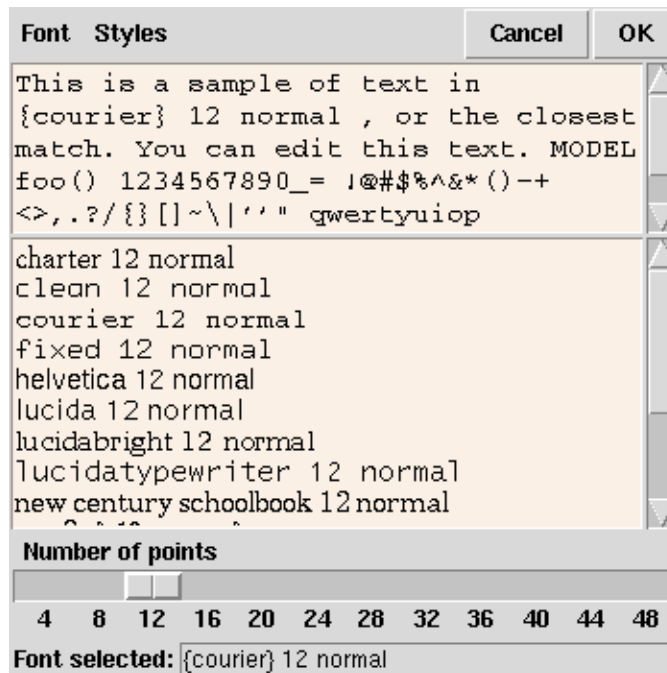


Figure 14-1 The font selection dialog.

To change the default fonts for ASCEND, see *Setting the Default Font* later in this chapter.

The font which the font selection dialog displays when it is opened is independent of the current window's font. This is actually a feature. When you close the font selection dialog (by pressing either the OK Button or the Cancel Button) and reopen it, it will display the same font as when it was closed. This way, once you find a font you like, you can change other ASCEND windows to this same font by simply opening the font selection dialog and pressing OK. As a default, the very first time you open the font dialog in an ASCEND session, the font is set to `Courier 12 normal`.

The font selection dialog has eight parts: Font Menu, Style Menu, Cancel Button, OK Button, Current Font Sample, Font Sampler Area, Point Size Slider, Current Font Selection.

## **14.2 FONT MENU**

The Font menu displays the fonts available for your platform (e.g., Helvetica, Courier). Selecting one of these fonts will update the Current Font Sample and Current Font Selection areas of the window.

## **14.3 STYLE MENU**

The Style menu allows you to specify attributes (e.g., Bold, Italic) for the selected font. As you add and remove attributes, the Current Font Sample and the Current Font Selection will reflect the changes.

## **14.4 CANCEL BUTTON**

The Cancel button closes the font selection window without changing the fonts of the window.

## **14.5 OK BUTTON**

The OK button closes the font selection window and sets the font of the window to the font listed in the Current Font Selection area.

## 14.6 CURRENT FONT SAMPLE

This area of the font selection window shows a sample of text in font, style, and size you have currently selected.

If you want to see what your current selection does to particular characters, you may type into this area. Note that your additions will be deleted when you change any aspect of the font (style, size, font).

## 14.7 FONT SAMPLER AREA

This area of the font selection window shows you a sample of the fonts available for your platform. You may make one of the listed fonts the current selection by clicking the font with the left mouse button. The currently selected styles and sizes remain in effect.

## 14.8 POINT SIZE SLIDER

This slider lets you choose the point size of the font. The text displayed in the Current Font Sample updates immediately.

## 14.9 CURRENT FONT SELECTION

This area displays the Tcl name for the font (including the size and style(s)) that you have currently selected. You may type in this area, but doing so will have no effect on the font.

## 14.10 SETTING THE DEFAULT FONT

To have ASCEND use the same font each time you run it, you need to do the following steps.

1. Use the font selection dialog to choose a font you like. Make a note of the Tcl name for the font; this name is displayed in the Current Font Selection area of the window.
2. Open the system utilities window and make a note of the value of ASCENDDIST.
3. Exit ASCEND.
4. Under the ASCENDDIST directory, there should be a directory

called TK, and in this directory a file called `ascend.ad`. Copy this file to your HOME directory<sup>1</sup> and name it `.ascend.ad`<sup>2</sup>.

5. Add the following lines at the end of `.ascend.ad`, replacing `courier 11 normal` with the font you noted in Step 1.

```
Global font courier 11 normal
Global labelfont courier 11 normal
Toolbox font courier 11 normal
Library font courier 11 normal
Display font courier 11 normal
Browser font courier 11 normal
Probe font courier 11 normal
Units font courier 11 normal
Script font courier 11 normal
Solver font courier 11 normal
Debugger font courier 11 normal
```

6. Save `.ascend.ad`, and restart ASCEND.

Note that this file also contains the default size and position for most ASCEND windows. To change the position or size of a window, edit the lines containing `geometry`; the format for the geometry is `WWxHH+xx+yy` where `WW` is the width of the window, `HH` is its height, `xx` is the distance between the left edge of the screen and the left edge of the window, and `yy` is the distance between the screen's top edge and the window's top edge.

- 
1. To set your HOME directory under Windows, open the Control Panel, double click the System icon, select the Environment tab, and set the HOME environment variable to a directory you want to consider "home".
  2. Under Windows, the name `_ascend.ad` also works.

# CHAPTER 15 THE PRINT DIALOG

## 15.1 OVERVIEW

The [print dialog](#) shown in [Figure 15-1](#) allows the user to modify the settings which control the printing of information from within ASCEND.

## 15.2 SETTINGS

### 15.2.1 DESTINATION

This is a pop-up menu that allows you to select one of the following options for printing: Print, Write to file, Append to file, Enscript, or Custom.



Figure 15-1 The print dialog.



### 15.2.1.1 PRINT

On UNIX machines, this option sends the window's contents to the printer specified in the Printer field (*PRINTER*). Under SystemV systems<sup>1</sup>, the command

```
lp -dPRINTER
```

is used as the interface to the printer; on all other UNIX systems, the command

```
lpr -PPRINTER
```

is used.

Under Windows, the command

```
notepad /p
```

is used to send the window's contents to the user's default printer.

### 15.2.1.2 WRITE TO FILE

Under UNIX, this option writes the contents of the window to the file listed in the Name of file field. If a file with the same name exists, ASCEND will overwrite the file after verifying that the user wants to overwrite the file.

This option is not available on the Windows platform.

This option is another version of Save As and will likely go away in future releases of ASCEND.

### 15.2.1.3 APPEND TO FILE

Under UNIX, this option appends the contents of the window to the file listed in the Name of file field. If the file does not exist, it will be created.

This option is not available on the Windows platform.

This option will likely go away in future releases of ASCEND.

### 15.2.1.4 ENSCRIPT

On UNIX, this option uses the `enscript` program to queue the window's contents to the printer specified in the Printer field (*PRINTER*). On SystemV systems, the command `enscript -dPRINTER enscript-flags` is used; on all other UNIX systems, the command

---

1. HP-UX, SGI IRIX, and Solaris 2.x.

`enscript -PPRINTER enscript-flags`

is used. The value of `enscript-flags` is the value specified in the Enscript flags field.

This option is not available on the Windows platform.

#### **15.2.1.5 CUSTOM**

This option allows the user to specify a custom print command. The user should type their custom command in the User print command field; the command should accept a file name as its final argument.

This option is not available on the Windows platform.

### **15.2.2 PRINTER**

Under UNIX, this field specifies the printer to send the document to when the Destination is Print or Enscript.

This field has no effect under Windows.

### **15.2.3 NAME OF FILE**

Under UNIX, this field contains the name of the file used by the Write to file and Append to file options.

This field has no effect under Windows.

### **15.2.4 ENSCRIPT FLAGS**

Under UNIX, this field contains the options sent to the `enscript` program when the Destination is Enscript.

This field has no effect under Windows.

### **15.2.5 USER PRINT COMMAND**

Under UNIX, this field contains the command used to “print” the window’s contents when the Destination is Custom. This command should accept the name of a file (a temporary file containing the contents of the window) as its final argument.

This field has no effect under Windows.

## **15.3 BUTTONS**

### **15.3.1 OK**

Pressing this button accepts the settings, sends the document to the printer or to the specified file, and closes the print dialog. The values displayed in the Text print command and the PRINTER fields in the system utilities window will change to reflect the new settings.

### **15.3.2 HELP**

Pressing this button should cause your web browser to display this document.

### **15.3.3 CANCEL**

This button ignores any changes you may have made to the settings and closes the print dialog. The file is not printed.

# CHAPTER 16 SOLVED SIMPLE MODELING PROBLEMS WITH ASCEND

In this chapter we present two simple modeling problems for which we then show you our ASCEND models for solving them. Modeling is a matter of style, and we will start to show you what we believe to be good styles for modeling. We assume you have not used the ASCEND system before. These problems are very generic and should be readily followed by anyone with a modest technical background.

One purpose is to show you some of the different ways you can use ASCEND. Specifically we want to show you that you can use ASCEND to setup and solve the simple types of problems that you might have solved using a spreadsheeting program. Indeed, we use ASCEND to solve homework problems quite often. When you factor in the powerful debugging tools, you might find it faster to use ASCEND, especially as the models get more complex. And no one would want (we think) to solve a 20,000 simultaneous nonlinear equation model using a spreadsheeting program.

A major advantage of using ASCEND is that once you have written, debugged and learned to solve such a model, you can interactively alter the "fixed" flags for the variables, changing which variables are to be fixed and which to be calculated. You can then immediately solve or optimize the new problem, using the previously solved problem as the initial guess.

## 16.1 ROOTS OF A POLYNOMIAL

In this problem you wish to find the roots of a polynomial. Assume you do not wish to keep the code. You could readily use a spreadsheet pro-

gram with its "root finder" routine to solve this type of problem, but you can as readily use ASCEND.

### 16.1.1 PROBLEM STATEMENT

Numerically compute the roots of  $(x-1)(x-5)(x+7)(x^2+1) = 0$ . (Given in this form the roots are obviously 1, 5, and -7. Two roots are complex, and ASCEND will not find them.)

### 16.1.2 ANSWER

You can find the roots by guessing initial points after typing in, loading and compiling the following model. You would use any text editor to enter this model into the computer. If you use a "WYSIWYG" (what you see is what you get) editor such as Word or Framemaker, be sure to save the file as a **text only** file. If possible, use a simpler text editor.

```
MODEL polynomial_roots;                                1
    x          IS_A generic_real;                       2
    (x-1)*(x-5)*(x+7)*(x^2+1) = 0;                     3
END polynomial_roots;                                  4
```

This simple model is a stand-alone model. You need no other predefined library models to support it. Load and compile an instance of this model (using tools in the LIBRARY tool set), browse it (using the BROWSER tool set) to see if it appears to have compiled correctly, and then pass it to the SOLVER tool set.

This model involves a single equation in the single unknown variable,  $x$ . The ASCEND solver treats a single equation in one unknown in a special manner when asked to solve it. The solver first attempts to rearrange the equation by simple algebraic manipulations to isolate the unknown on the left hand side of the equation in the form  $x =$  expression not involving  $x$ . In this form, solving is simply evaluating the expression on the right hand side once. Here the solver would fail as there is no way to isolate  $x$  on the left hand side as the equation is a fifth order polynomial in  $x$ . When rearrangement fails, the solver uses bisection to locate a root in the range between the lower and upper bound on the variable. You can see the bounds,  $x.lower$  and  $x.upper$ , using the BROWSER. The default values for these bounds are plus and minus  $10^{20}$  respectively, which gives a very large range in which to look for the root. You should change the bounds<sup>1</sup> to more realistic

---

1. To set the value for a variable interactively, select the variable when it is display in the right window or in the lower window with the RIGHT mouse button (the other button). A window for changing its value opens.

ones. Selecting bounds to be -10 to 0 and then solving will find the root  $x=-7$ . Selecting other values will find the other roots.

This model illustrates that you can quickly set up and solve simple problems using ASCEND. Note that you would have been required to place bounds on  $x$  had you used a goal seeking tool in a spreadsheeting program if you wanted to control which root to locate.

## 16.2 NUMERICAL INTEGRATION OF TABULAR DATA

This problem is similar to the previous one in that it is very easy to set up and solve. It adds in the notion of units (e.g., ft, m, hr, atm) which ASCEND handles in a straight-forward manner, relieving modelers from thinking about converting among the many units they might use when expressing the data for a problem.

Again we are talking about producing throw-away code. All we are really concerned with here is the answer which we intend to put into a report. We are using ASCEND as a "calculator."

### 16.2.1 PROBLEM STATEMENT

Given the following velocity data vs. time, estimate numerically the distance one has traveled between time equal to zero and 100 seconds.

**Table 1: Velocity data to be integrated**

<b>data point number</b>	<b>time, s</b>	<b>velocity, ft/min</b>
1	0	100
2	10	120
3	20	130
4	30	135
5	40	140
6	50	160
7	60	180
8	70	210
9	80	240
10	90	220
11	100	200

## 16.2.2 ANSWER

(This example will be solved using variables whose types are defined in the file *atoms.lib*. You must load this file first before loading the file with the code below, else you will experience a number of diagnostic messages indicating missing type definitions. See the document entitled (\*\*\*\*link\*\*\*\* *library\_example.fm5*, \*\*\*\*link label\*\*\*\* the ASCEND predefined collection of models) for a discussion of libraries on ASCEND.)

The distance traveled is the integral of the velocity over time. We can use Simpson's rule to carry out this integration for evenly spaced points.

$$d = ((v[1] + 4 v[2] + v[3]) + (v[3] + 4 v[4] + v[5]) + \dots + (v[N-2] + 4 v[N-1] + v[N])) * \Delta t / 6 \quad (16.1)$$

where  $d$  is the distance covered when traveling at the velocities,  $V[k]$ , listed. This formula requires there to be an odd number of 3 or more evenly spaced data points, which is fine here as we have eleven velocity points evenly spaced in time. (If there had been an even number of points, we could use Simpson's rule for all but the last time interval and use a simple trapezoidal rule to integrate it.)

An ASCEND model to evaluate this distance is as follows. The types definitions for *speed*, *time* and *distance* are in the file *atoms.lib*.

```

MODEL travel_distance;                                1
    kmax          IS_A integer_constant;              2
    v[1..2*kmax+1] IS_A speed;                        3
    delta_time    IS_A time;                          4
    d             IS_A distance;                      5
                                                        6
    d = SUM[v[2*k-1]+4*v[2*k]+v[2*k+1] SUCH_THAT k IN
            [1..kmax]]*delta_time/6;                  7
END travel_distance;                                  8
                                                        9

MODEL test_travel_distance REFINES travel_distance;   10
    kmax          ::= 5;                               11
                                                        12

METHODS                                               13
    METHOD specify;                                    14
        v[1..2*kmax+1].fixed := TRUE;                15
        delta_time.fixed      := TRUE;                16
    END specify;                                       17
                                                        18

    METHOD values;                                     19
        v[1]                  := 100 {ft/min};       20

```

```

v[2] := 120 {ft/min}; 21
v[3] := 130 {ft/min}; 22
v[4] := 135 {ft/min}; 23
v[5] := 140 {ft/min}; 24
v[6] := 160 {ft/min}; 25
v[7] := 180 {ft/min}; 26
v[8] := 210 {ft/min}; 27
v[9] := 240 {ft/min}; 28
v[10] := 220 {ft/min}; 29
v[11] := 200 {ft/min}; 30
delta_time := 10 {s}; 31
END values; 32
END test_travel_distance; 33
```

If you look carefully at this model, you will note that we did NOT account for the conversion factors required because velocities are in ft/min while the time increment is in seconds. ASCEND understands these units and makes all the needed conversions. When you run this model, you can ask for the distance to be displayed to you in any supported length units you would prefer (e.g., ft, mile, m, cm, angstroms, light-years). The distance traveled, when reported using SI units, is 42.84 m.





## CHAPTER 17 A CONDITIONAL MODELING EXAMPLE: REPRESENTING A SUPERSTRUCTURE

To give an example of the application of the conditional modeling tool in ASCEND -the WHEN statement-, we developed a simplified model for the superstructure given in Figure 17-1. The code listed below exists in a file in the ASCEND models subdirectory entitled *when\_demo.a4c*. You could run this example by loading this file and using it and its corresponding script *when\_demo.a4s*.

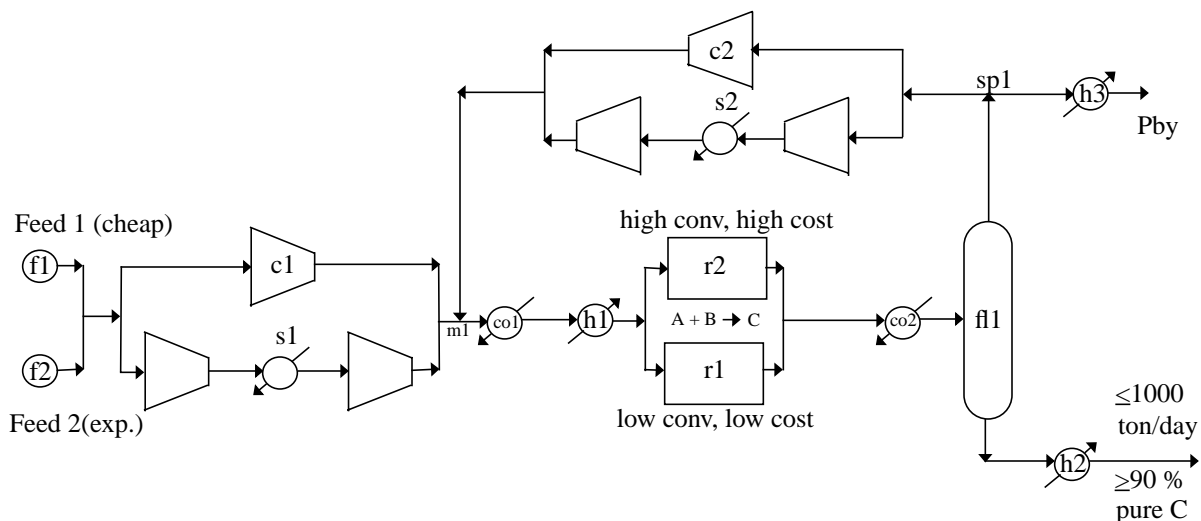


Figure 17-1 Superstructure used in the example of the application of the when statement

### 17.1 THE WHEN STATEMENT

Before showing the example, we want to start by giving a brief explanation about the semantics of the WHEN statement, a tool which allows ASCEND to represent conditional models efficiently.

In the WHEN statement, we take advantage of the fact that ASCEND is based on object oriented concepts where model definitions can contain parts that contain parts to any level. Furthermore, in ASCEND, a simple

relation is treated as an object by itself and can have a name. Based on these ideas, the syntax for the WHEN statement is:

```

WHEN (list_of_variables)
    CASE list_of_values_1:
        USE name_of_equation_1;
        USE name_of_model_1;
    CASE list_of_values_2:
        USE name_of_equation_2;
        USE name_of_model_2;
    CASE list_of_values_nminus1:
        USE
name_of_equation_nminus1;
        USE name_of_model_nminus1;
    OTHERWISE:
        USE name_of_equation_n;
        USE name_of_model_n;
END;

```

The following are important observations about the implementation:

- 1 The WHEN statement does not mean conditional compilation. We create and have available the data structures for all of the variables and equations in each of the models. This is actually a requirement for the solution algorithms of conditional models. All the models and equations whose name is given in each of the cases should be declared inside the model which contains the WHEN statement.
- 2 The variables in the list of variables can be of any type among boolean, integer or symbol or any combination of them. That is, we are not limited to the use of boolean variables. Obviously, The list of values in each case must be in agreement with the list of variables in the number of elements and type of each element. In other words, order matters in the list of variables of the WHEN statement, and parentheses are enclosing this list to make clear such a feature.
- 3 Names of arrays of models or equations are also allowed inside the scope of each CASE.

The WHEN statement represents an important contribution to modeling: it allows the user to define the domain of validity of both *models* and *equations* inside the cases of a WHEN statement. This feature enormously increases the scope of modeling in an equation based modeling environment.

Mainly, there are two different ways in which the WHEN statement can be used.:

- First, the WHEN statement can be used to select a configuration

of a problem among several alternative configurations.

- Second, in combination with logical relations, the WHEN statement can be used for conditional programming. That is, a problem in which the system of equations to be solved depends on the solution of the problem. A typical example of this situation is the laminar-turbulent flow transition. The selection of the equation to calculate the friction factor depends on the value of the Reynolds number, which is an unknown in the problem.

## 17.2 THE PROBLEM DESCRIPTION

In the example, there are two alternative feedstocks, two possible choices of the reactor and two choices for each of the compression systems. The user has to make 4 decisions (for example, using either the cheap feed or the expensive feed), therefore, there are  $2^4 = 16$  feasible configurations of the problem. All these 16 configurations are encapsulated in one ASCEND model containing 4 WHEN statements which depend on the value of 4 boolean variables.

The value of the four boolean variables will determine the structure of the problem to be solved. In this example, those values are defined by the modeler, but they also could be defined by some logic inference algorithm which would allow the automatic change of the structure of the problem.

The following section gives the code for this model. The first models correspond to the different types of unit operations existing in the superstructure. Those model are very simplified. You may want to skip them and analyze only the model *flowsheet*, in which the use and syntax of the WHEN statement as well as the configuration of the superstructure become evident.

## 17.3 THE CODE

As the code is in our ASCEND examples subdirectory, it has header information that we required of all such files included as one large comment extending over several lines. Comments are in the form (\* comment \*). The last item in this header information is a list of the files one must load before loading this one, i.e., *system.lib* and *atoms.lib*.

```
( *****\
                                     34
                                     when_demo.a4c
                                     35
```

```

by Vicente Rico-Ramirez                                     36
Part of the Ascend Library                                 37
                                                         38
This file is part of the Ascend modeling library.         39
The Ascend modeling library is free software; you can redistribute 40
it and/or modify it under the terms of the GNU General Public License as 41
published by the Free Software Foundation; either version 2 of the      42
License, or (at your option) any later version.          43
                                                         44
The Ascend Language Interpreter is distributed in hope that it will be 45
useful, but WITHOUT ANY WARRANTY; without even the implied warranty of 46
MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE. See the GNU       47
General Public License for more details.                  48
                                                         49

You should have received a copy of the GNU General Public License along with50
the program; if not, write to the Free Software Foundation, Inc., 675    51
Mass Ave, Cambridge, MA 02139 USA. Check the file named COPYING.       52
                                                         53

Use of this module is demonstrated by the associated script file        54
when_demo.s.                                                 55
\*****                                                    56
(*****\                                                    57
  $Date: 97/06/03 12:22:29 $                                     58
  $Revision: 1.1 $                                             59
  $Author: rv2a $                                             60
  $Source:/afs/cs.cmu.edu/project/ascend/Repository/models/when_demo.a4c,v $61
\*****                                                    62
                                                         63
(*)                                                           64
This model is intended to demonstrate the degree of flexibility         65
that the use of conditional statements -when statements- provides        66
to the representation of superstructures. We hope that this            67
application will become clear by looking at the MODEL flowsheet,       68
in which the existence/nonexistence of some of the unit operations      69
is represented by WHEN statements. A particular combination of         70
user defined boolean variables -see the methods values, configuration2, and 71
configuration3- will a define a particular configuration of the         72
problem.                                                       73
                                                         74

This model requires:                                           75
    "system.lib"                                             76
    "atoms.lib"                                             77
*)                                                           78
(* ***** *)                                               79
                                                         80

MODEL mixture;                                               81

```

```

components          IS_A set OF symbol_constant;      82
Cpi[components]    IS_A heat_capacity;          83
y[components]      IS_A fraction;              84
P                  IS_A pressure;              85
T                  IS_A temperature;           86
Cp                 IS_A heat_capacity;         87
                                                            88
                                                            89
SUM[y[i] | i IN components] = 1.0;            90
Cp = SUM[Cpi[i] * y[i] | i IN components];    91
                                                            92
METHODS                                                  93
  METHOD clear;                                          94
    y[components].fixed := FALSE;                    95
    Cpi[components].fixed := FALSE;                  96
    Cp.fixed := FALSE;                                97
    P.fixed := FALSE;                                 98
    T.fixed := FALSE;                                 99
  END clear;                                           100
                                                            101
  METHOD specify;                                       102
    Cpi[components].fixed := TRUE;                   103
    P.fixed := TRUE;                                  104
    T.fixed := TRUE;                                  105
    y[components].fixed := TRUE;                     106
    y[CHOICE[components]].fixed := FALSE;            107
  END specify;                                         108
                                                            109
  METHOD reset;                                         110
    RUN clear;                                        111
    RUN specify;                                      112
  END reset;                                           113
END mixture;                                           114
                                                            115
(* ***** *)                                       116
                                                            117
MODEL molar_stream;                                    118
  state          IS_A mixture;                        119
  Ftot,f[components] IS_A molar_rate;                120
  components     IS_A set OF symbol_constant;        121
  P              IS_A pressure;                       122
  T              IS_A temperature;                    123
  Cp            IS_A heat_capacity;                   124
                                                            125
  components, state.components ARE_THE_SAME;          126
  P, state.P ARE_THE_SAME;                            127

```

```

T, state.T                ARE_THE_SAME;                128
Cp, state.Cp              ARE_THE_SAME;                129
                                                                    130
FOR i IN components CREATE 131
    f_def[i]: f[i] = Ftot*state.y[i];                132
END;                                                       133
                                                                    134
METHODS                                                     135
    METHOD clear;                                          136
        RUN state.clear;                                137
        Ftot.fixed:= FALSE;                             138
        f[components].fixed:= FALSE;                   139
    END clear;                                           140
                                                                    141
    METHOD specify;                                       142
        RUN state.specify;                              143
        state.y[components].fixed := FALSE;           144
        f[components].fixed := TRUE;                  145
    END specify;                                         146
                                                                    147
    METHOD reset;                                         148
        RUN clear;                                     149
        RUN specify;                                   150
    END reset;                                           151
END molar_stream;                                        152
(* ***** *)                                         153
                                                                    154
MODEL cheap_feed;                                        155
    stream          IS_A molar_stream;                 156
    cost_factor     IS_A cost_per_mole;                157
    cost            IS_A cost_per_time;                158
                                                                    159
    stream.f['A'] = 0.060 {kg_mole/s};                 160
    stream.f['B'] = 0.025 {kg_mole/s};                 161
    stream.f['D'] = 0.015 {kg_mole/s};                 162
    stream.f['C'] = 0.00 {kg_mole/s};                 163
    stream.T = 300 {K};                                164
    stream.P = 5 {bar};                                165
                                                                    166
    cost = cost_factor * stream.Ftot;                   167
METHODS                                                     168
    METHOD clear;                                          169
        RUN stream.clear;                              170
        cost_factor.fixed := FALSE;                   171
        cost.fixed := FALSE;                           172
    END clear;                                           173

```

```

METHOD specify;
  RUN stream.specify;
  stream.f[stream.components].fixed := FALSE;
  cost_factor.fixed := TRUE;
  stream.T.fixed := FALSE;
  stream.P.fixed := FALSE;
END specify;

METHOD reset;
  RUN clear;
  RUN specify;
END reset;

END cheap_feed;

(* ***** *)

MODEL expensive_feed;
  stream          IS_A molar_stream;
  cost_factor     IS_A cost_per_mole;
  cost            IS_A cost_per_time;

  stream.f['A'] = 0.065 {kg_mole/s};
  stream.f['B'] = 0.030 {kg_mole/s};
  stream.f['D'] = 0.05  {kg_mole/s};
  stream.f['C'] = 0.00  {kg_mole/s};
  stream.T = 320 {K};
  stream.P = 6 {bar};

  cost = 3 * cost_factor * stream.Ftot;

METHODS
  METHOD clear;
    RUN stream.clear;
    cost_factor.fixed := FALSE;
    cost.fixed := FALSE;
  END clear;

  METHOD specify;
    RUN stream.specify;
    stream.f[stream.components].fixed := FALSE;
    cost_factor.fixed := TRUE;
    stream.T.fixed := FALSE;
    stream.P.fixed := FALSE;
  END specify;

```



```

METHOD reset;
  RUN clear;
  RUN specify;
END reset;

END expensive_feed;

(* ***** *)

MODEL heater;
  input,output      IS_A molar_stream;
  heat_supplied     IS_A energy_rate;
  components        IS_A set OF symbol_constant;
  cost              IS_A cost_per_time;
  cost_factor       IS_A cost_per_energy;

  components,input.components,output.components ARE_THE_SAME;

  input.state.Cpi[components],
    output.state.Cpi[components] ARE_THE_SAME;

  FOR i IN components CREATE
    input.f[i] = output.f[i];
  END;

  input.P = output.P;

  heat_supplied = input.Cp *(output.T - input.T) * input.Ftot;

  cost = cost_factor * heat_supplied;

METHODS
  METHOD clear;
    RUN input.clear;
    RUN output.clear;
    cost.fixed := FALSE;
    cost_factor.fixed := FALSE;
    heat_supplied.fixed := FALSE;
  END clear;

  METHOD specify;
    RUN input.specify;
    cost_factor.fixed := TRUE;

```

```

        heat_supplied.fixed := TRUE;                266
    END specify;                                    267
                                                    268
    METHOD seqmod;                                    269
        cost_factor.fixed := TRUE;                  270
        heat_supplied.fixed := TRUE;                271
    END seqmod;                                      272
                                                    273
    METHOD reset;                                     274
        RUN clear;                                  275
        UN specify;                                  276
    END reset;                                       277
                                                    278
END heater;                                         279
                                                    280
                                                    281
                                                    282
(* ***** *)                                     283
                                                    284
                                                    285
MODEL cooler;                                       286
                                                    287
    input,output      IS_A molar_stream;            288
    heat_removed      IS_A energy_rate;            289
    components        IS_A set OF symbol_constant; 290
    cost              IS_A cost_per_time;          291
    cost_factor       IS_A cost_per_energy;        292
                                                    293
    components,input.components,output.components  ARE_THE_SAME; 294
    input.state.Cpi[components],                  295
        output.state.Cpi[components]              ARE_THE_SAME; 296
                                                    297
    FOR i IN components CREATE                      298
        input.f[i] = output.f[i];                 299
    END;                                            300
    input.P = output.P;                            301
    heat_removed = input.Cp *(input.T - output.T) * input.Ftot; 302
                                                    303
    cost = cost_factor * heat_removed;             304
                                                    305
METHODS                                             306
    METHOD clear;                                    307
        RUN input.clear;                            308
        RUN output.clear;                          309
        cost.fixed := FALSE;                        310
        cost_factor.fixed := FALSE;                 311

```

```

    heat_removed.fixed := FALSE;                                312
END clear;                                                    313
                                                            314
METHOD specify;                                              315
    RUN input.specify;                                       316
    cost_factor.fixed := TRUE;                                317
    heat_removed.fixed := TRUE;                               318
END specify;                                                319
                                                            320
METHOD seqmod;                                              321
    cost_factor.fixed := TRUE;                                322
    heat_removed.fixed := TRUE;                               323
END seqmod;                                                 324
                                                            325
METHOD reset;                                               326
    RUN clear;                                               327
    RUN specify;                                             328
END reset;                                                  329
                                                            330
END cooler;                                                 331
                                                            332
                                                            333
(* ***** *)                                             334
                                                            335
                                                            336
MODEL single_compressor; (* Adiabatic Compression *)        337
                                                            338
    input,output      IS_A molar_stream;                     339
    components        IS_A set OF symbol_constant;          340
    work_supplied     IS_A energy_rate;                       341
    gamma             IS_A factor;                            342
    pressure_rate     IS_A factor;                            343
    R                 IS_A gas_constant;                     344
    cost              IS_A cost_per_time;                    345
    cost_factor       IS_A cost_per_energy;                  346
                                                            347
    components,input.components,output.components ARE_THE_SAME; 348
    input.state.Cpi[components],                    349
        output.state.Cpi[components] ARE_THE_SAME;          350
                                                            351
    FOR i IN components CREATE                             352
        input.f[i] = output.f[i];                          353
    END;                                                  354
                                                            355
    gamma = input.Cp / (input.Cp - R);                     356
    pressure_rate = output.P / input.P;                     357

```

```

output.T = input.T * (pressure_rate ^((gamma-1)/gamma) );           358
                                                                    359
work_supplied = input.Ftot * gamma * R *input.T                    360
    * (gamma/(gamma -1))                                           361
    * (( (pressure_rate)^((gamma -1)/gamma))-1 );                  362
                                                                    363
cost = cost_factor * work_supplied;                                 364
                                                                    365
METHODS                                                             366
METHOD clear;                                                       367
    RUN input.clear;                                                368
    RUN output.clear;                                              369
    gamma.fixed := FALSE;                                          370
    cost.fixed := FALSE;                                           371
    cost_factor.fixed := FALSE;                                     372
    pressure_rate.fixed := FALSE;                                   373
    work_supplied.fixed := FALSE;                                   374
END clear;                                                          375
                                                                    376
METHOD specify;                                                     377
    RUN input.specify;                                             378
    pressure_rate.fixed := TRUE;                                    379
END specify;                                                        380
                                                                    381
METHOD seqmod;                                                      382
    cost_factor.fixed := TRUE;                                      383
    pressure_rate.fixed := TRUE;                                    384
END seqmod;                                                         385
                                                                    386
METHOD reset;                                                       387
    RUN clear;                                                     388
    RUN specify;                                                   389
END reset;                                                          390
END single_compressor;                                             391
(* ***** *)                                                    392
                                                                    393
MODEL staged_compressor;                                           394
                                                                    395
    input,output           IS_A molar_stream;                       396
    components             IS_A set OF symbol_constant;            397
    work_supplied          IS_A energy_rate;                        398
    T_middle               IS_A temperature;                       399
    heat_removed           IS_A energy_rate;                       400
    gamma                  IS_A factor;                            401
    pressure_rate          IS_A factor;                             402
    R                      IS_A gas_constant;                      403

```

```

cost          IS_A cost_per_time;          404
cost_factor_work IS_A cost_per_energy;    405
cost_factor_heat IS_A cost_per_energy;    406
                                                    407
components,input.components,output.components ARE_THE_SAME; 408
input.state.Cpi[components],                409
    output.state.Cpi[components]            ARE_THE_SAME; 410
                                                    411
FOR i IN components CREATE                  412
    input.f[i] = output.f[i];              413
END;                                        414
                                                    415
gamma = input.Cp / (input.Cp - R);         416
output.T = input.T;                        417
pressure_rate = output.P / input.P;       418
T_middle = input.T * (pressure_rate ^((gamma-1)/gamma)); 419
                                                    420
work_supplied = input.Ftot * 2 * gamma * R *input.T
    * (gamma/(gamma-1))                    422
    * (( (pressure_rate)^((gamma -1)/( 2*gamma)))-1 ); 423
                                                    424
heat_removed = input.Ftot * input.Cp * (T_middle - input.T); 425
                                                    426
cost = cost_factor_work * work_supplied + 427
    cost_factor_heat * heat_removed;       428
                                                    429
METHODS                                     430
METHOD clear;                               431
    RUN input.clear;                        432
    RUN output.clear;                      433
    gamma.fixed := FALSE;                  434
    pressure_rate.fixed := FALSE;          435
    T_middle.fixed := FALSE;              436
    work_supplied.fixed := FALSE;          437
    heat_removed.fixed := FALSE;           438
    cost_factor_heat.fixed := FALSE;       439
    cost_factor_work.fixed := FALSE;       440
    cost.fixed := FALSE;                   441
END clear;                                  442
                                                    443
METHOD specify;                             444
    RUN input.specify;                     445
    cost_factor_heat.fixed := TRUE;        446
    cost_factor_work.fixed := TRUE;        447
END specify;                                448
                                                    449

```

```

METHOD seqmod;                                450
    cost_factor_heat.fixed := TRUE;           451
    cost_factor_work.fixed := TRUE;           452
    pressure_rate.fixed := TRUE;              453
END seqmod;                                    454
                                              455

METHOD reset;                                  456
    RUN clear;                                457
    RUN specify;                              458
END reset;                                    459
END staged_compressor;                        460
                                              461

(* ***** *)                                462
                                              463

MODEL mixer;                                  464
                                              465

    components          IS_A set OF symbol_constant; 466
    n_inputs            IS_A integer_constant;        467
    feed[1..n_inputs], out IS_A molar_stream;         468
    To                  IS_A temperature;             469
                                              470

    components,feed[1..n_inputs].components,        471
    out.components          ARE_THE_SAME;             472
                                              473

    feed[1..n_inputs].state.Cpi[components],        474
    out.state.Cpi[components] ARE_THE_SAME;          475
                                              476

    FOR i IN components CREATE                     477
        cmb[i]: out.f[i] = SUM[feed[1..n_inputs].f[i]]; 478
    END;                                             479
                                              480

    SUM[(feed[i].Cp *feed[i].Ftot * (feed[i].T - To))|i IN [1..n_inputs]]= 481
        out.Cp *out.Ftot * (out.T - To);           482
                                              483

    SUM[( feed[i].Ftot * feed[i].T / feed[i].P )|i IN [1..n_inputs]] = 484
        out.Ftot * out.T / out.P;                 485
                                              486

METHODS                                         487
    METHOD clear;                                 488
        RUN feed[1..n_inputs].clear;             489
        RUN out.clear;                           490
        To.fixed := FALSE;                       491
    END clear;                                   492
                                              493

    METHOD specify;                               494
        To.fixed := TRUE;                        495

```

```

    RUN feed[1..n_inputs].specify;           496
END specify;                               497
                                           498
METHOD seqmod;                             499
    To.fixed := TRUE;                       500
END seqmod;                                501
                                           502
METHOD reset;                              503
    RUN clear;                             504
    RUN specify;                           505
END reset;                                 506
                                           507
END mixer;                                 508
                                           509
(* ***** *)                             510
                                           511
MODEL splitter;                            512
    components          IS_A set OF symbol_constant; 513
    n_outputs           IS_A integer_constant;      514
    feed, out[1..n_outputs] IS_A molar_stream;     515
    split[1..n_outputs] IS_A fraction;             516
                                           517
    components, feed.components,              518
    out[1..n_outputs].components ARE_THE_SAME;    519
                                           520
    feed.state,                               521
    out[1..n_outputs].state ARE_THE_SAME;        522
                                           523
    FOR j IN [1..n_outputs] CREATE           524
        out[j].Ftot = split[j]*feed.Ftot;      525
    END;                                       526
                                           527
    SUM[split[1..n_outputs]] = 1.0;          528
                                           529
METHODS                                     530
    METHOD clear;                             531
        RUN feed.clear;                     532
        RUN out[1..n_outputs].clear;        533
        split[1..n_outputs-1].fixed :=FALSE; 534
    END clear;                               535
                                           536
    METHOD specify;                           537
        RUN feed.specify;                   538
        split[1..n_outputs-1].fixed:=TRUE;  539
    END specify;                             540
                                           541

```

```

METHOD seqmod;
  split[1..n_outputs-1].fixed:=TRUE;
END seqmod;

METHOD reset;
  RUN clear;
  RUN specify;
END reset;
END splitter;

(* ***** *)

MODEL cheap_reactor;
  components          IS_A set OF symbol_constant;
  input, output       IS_A molar_stream;
  low_turnover        IS_A molar_rate;
  stoich_coef[input.components] IS_A factor;
  cost_factor         IS_A cost_per_mole;
  cost                IS_A cost_per_time;

  components,input.components, output.components ARE_THE_SAME;
  input.state.Cpi[components],
  output.state.Cpi[components] ARE_THE_SAME;

  FOR i IN input.components CREATE
    output.f[i] = input.f[i] + stoich_coef[i]*low_turnover;
  END;

  input.T = output.T;
  (* ideal gas constant volume *)
  input.Ftot * input.T / input.P = output.Ftot * output.T/output.P;

  cost = cost_factor * low_turnover;

METHODS
  METHOD clear;
    RUN input.clear;
    RUN output.clear;
    low_turnover.fixed:= FALSE;
    stoich_coef[input.components].fixed:= FALSE;
    cost.fixed := FALSE;
    cost_factor.fixed := FALSE;
  END clear;

  METHOD specify;
    RUN input.specify;

```



```

    low_turnover.fixed:= TRUE;                               588
    stoich_coef[input.components].fixed:= TRUE;             589
    cost_factor.fixed := TRUE;                              590
END specify;                                               591
                                                            592
METHOD seqmod;                                             593
    low_turnover.fixed:= TRUE;                               594
    stoich_coef[input.components].fixed:= TRUE;             595
    cost_factor.fixed := TRUE;                              596
END seqmod;                                                597
                                                            598
METHOD reset;                                              599
    RUN clear;                                              600
    RUN specify;                                            601
END reset;                                                 602
                                                            603
END cheap_reactor;                                        604
                                                            605
(* ***** *)                                           606
                                                            607
MODEL expensive_reactor;                                   608
                                                            609
    components          IS_A set OF symbol_constant;       610
    input, output       IS_A molar_stream;                 611
    high_turnover        IS_A molar_rate;                  612
    stoich_coef[input.components] IS_A factor;             613
    cost_factor          IS_A cost_per_mole;               614
    cost                 IS_A cost_per_time;               615
                                                            616
    components,input.components, output.components ARE_THE_SAME; 617
                                                            618
    input.state.Cpi[components],                          619
    output.state.Cpi[components] ARE_THE_SAME;            620
                                                            621
    FOR i IN input.components CREATE                       622
        output.f[i] = input.f[i] + stoich_coef[i]*high_turnover; 623
    END;                                                  624
                                                            625
    input.T = output.T;                                    626
    (* ideal gas constant volume *)                       627
    input.Ftot * input.T / input.P = output.Ftot * output.T/output.P; 628
                                                            629
    cost = cost_factor * high_turnover;                   630
                                                            631
METHODS                                                    632
                                                            633

```

```

METHOD clear;                                634
  RUN input.clear;                            635
  RUN output.clear;                          636
  high_turnover.fixed:= FALSE;               637
  stoich_coef[input.components].fixed:= FALSE; 638
  cost.fixed := FALSE;                       639
  cost_factor.fixed := FALSE;               640
END clear;                                    641
                                              642

METHOD specify;                              643
  RUN input.specify;                         644
  high_turnover.fixed:= TRUE;               645
  stoich_coef[input.components].fixed:= TRUE; 646
  cost_factor.fixed := TRUE;               647
END specify;                                  648
                                              649

METHOD seqmod;                               650
  high_turnover.fixed:= TRUE;               651
  stoich_coef[input.components].fixed:= TRUE; 652
  cost_factor.fixed := TRUE;               653
END seqmod;                                   654
                                              655

METHOD reset;                               656
  RUN clear;                                 657
  RUN specify;                              658
END reset;                                    659
                                              660

END expensive_reactor;                       661
                                              662

(* ***** *)                              663
                                              664

MODEL flash;                                 665
  components          IS_A set OF symbol_constant; 666
  feed,vap,liq        IS_A molar_stream;          667
  alpha[feed.components] IS_A factor;             668
  ave_alpha           IS_A factor;               669
  vap_to_feed_ratio   IS_A fraction;             670
                                              671

  components,feed.components,                672
  vap.components,                             673
  liq.components          ARE_THE_SAME;         674
                                              675

  feed.state.Cpi[components],                676
  vap.state.Cpi[components],                 677
  liq.state.Cpi[components] ARE_THE_SAME;     678
                                              679

```

```

vap_to_feed_ratio*feed.Ftot = vap.Ftot;                                680
                                                                    681
FOR i IN feed.components CREATE                                       682
    cmb[i]: feed.f[i] = vap.f[i] + liq.f[i];                          683
    eq[i]: vap.state.y[i]*ave_alpha = alpha[i]*liq.state.y[i];      684
END;                                                                    685
                                                                    686
feed.T = vap.T;                                                       687
feed.T = liq.T;                                                       688
feed.P = vap.P;                                                       689
feed.P = liq.P;                                                       690
                                                                    691
METHODS                                                                  692
                                                                    693
METHOD clear;                                                          694
    RUN feed.clear;                                                    695
    RUN vap.clear;                                                     696
    RUN liq.clear;                                                     697
    alpha[feed.components].fixed:=FALSE;                              698
    ave_alpha.fixed:=FALSE;                                            699
    vap_to_feed_ratio.fixed:=FALSE;                                    700
END clear;                                                             701
                                                                    702
METHOD specify;                                                       703
    RUN feed.specify;                                                  704
    alpha[feed.components].fixed:=TRUE;                                705
    vap_to_feed_ratio.fixed:=TRUE;                                     706
END specify;                                                           707
                                                                    708
METHOD seqmod;                                                         709
    alpha[feed.components].fixed:=TRUE;                                710
    vap_to_feed_ratio.fixed:=TRUE;                                     711
END seqmod;                                                            712
                                                                    713
METHOD reset;                                                         714
    RUN clear;                                                         715
    RUN specify;                                                       716
END reset;                                                             717
END flash;                                                             718
                                                                    719

```

Next, the model *flowsheet* is presented. This model represents one of the applications of the WHEN statement. Namely, selecting among alternative configurations of the problem. Note that in each of the WHEN statements we define the conditional existence of complete ASCEND models. A specific combination for each of the conditional

variables -boolean\_vars in the example- will define a specific configuration of the problem. Once a configuration has been selected, it will be kept until the user decides to change it. Note that the user does not have to recompile the model to switch among alternative configurations. The reconfiguration of the system can be done automatically by simply changing the values of the conditional variables. An obvious application of this would be the synthesis of process networks. While running the script *when\_demo.a4s*, note the changes in the number of active equations, active variables and fixed variables for the different configurations. For example, the configuration defined by one of the feeds, two single-stage compressors and one of the reactors contains 169 active equations.

```
( * ***** *) 720
721
722
MODEL flowsheet; 723
724
(* units *) 725
726
f1      IS_A cheap_feed; 727
f2      IS_A expensive_feed; 728
729
c1      IS_A single_compressor; 730
s1      IS_A staged_compressor; 731
732
c2      IS_A single_compressor; 733
s2      IS_A staged_compressor; 734
735
r1      IS_A cheap_reactor; 736
r2      IS_A expensive_reactor; 737
738
col1,co2 IS_A cooler; 739
h1,h2,h3 IS_A heater; 740
fl1     IS_A flash; 741
sp1     IS_A splitter; 742
m1     IS_A mixer; 743
744
(* boolean variables *) 745
746
select_feed1 IS_A boolean_var; 747
select_single1 IS_A boolean_var; 748
select_cheapr1 IS_A boolean_var; 749
select_single2 IS_A boolean_var; 750
751
(* define sets *) 752
```

		753
m1.n_inputs :=2;		754
sp1.n_outputs := 2;		755
		756
(* wire up flowsheet *)		757
		758
f1.stream, f2.stream, c1.input, s1.input	ARE_THE_SAME;	759
c1.output, s1.output, m1.feed[2]	ARE_THE_SAME;	760
m1.out, col.input	ARE_THE_SAME;	761
col.output, h1.input	ARE_THE_SAME;	762
h1.output, r1.input, r2.input	ARE_THE_SAME;	763
r1.output, r2.output, co2.input	ARE_THE_SAME;	764
co2.output, fl1.feed	ARE_THE_SAME;	765
fl1.liq, h2.input	ARE_THE_SAME;	766
fl1.vap, sp1.feed	ARE_THE_SAME;	767
sp1.out[1], h3.input	ARE_THE_SAME;	768
sp1.out[2], c2.input, s2.input	ARE_THE_SAME;	769
c2.output, s2.output, m1.feed[1]	ARE_THE_SAME;	770
		771
		772
(* Conditional statements *)		773
		774
WHEN (select_feed1)		775
CASE TRUE:		776
USE f1;		777
CASE FALSE:		778
USE f2;		779
END;		780
		781
WHEN (select_single1)		782
CASE TRUE:		783
USE c1;		784
CASE FALSE:		785
USE s1;		786
END;		787
		788
WHEN (select_cheapr1)		789
CASE TRUE:		790
USE r1;		791
CASE FALSE:		792
USE r2;		793
END;		794
		795
WHEN (select_single2)		796
CASE TRUE:		797
USE c2;		798

```
CASE FALSE: 799
  USE s2; 800
END; 801
802
803
METHODS 804
  METHOD clear; 805
    RUN f1.clear; 806
    RUN f2.clear; 807
    RUN c1.clear; 808
    RUN c2.clear; 809
    RUN s1.clear; 810
    RUN s2.clear; 811
    RUN co1.clear; 812
    RUN co2.clear; 813
    RUN h1.clear; 814
    RUN h2.clear; 815
    RUN h3.clear; 816
    RUN r1.clear; 817
    RUN r2.clear; 818
    RUN fl1.clear; 819
    RUN sp1.clear; 820
    RUN m1.clear; 821
  END clear; 822
823
  METHOD seqmod; 824
    RUN c1.seqmod; 825
    RUN c2.seqmod; 826
    RUN s1.seqmod; 827
    RUN s2.seqmod; 828
    RUN co1.seqmod; 829
    RUN co2.seqmod; 830
    RUN h1.seqmod; 831
    RUN h2.seqmod; 832
    RUN h3.seqmod; 833
    RUN r1.seqmod; 834
    RUN r2.seqmod; 835
    RUN fl1.seqmod; 836
    RUN sp1.seqmod; 837
    RUN m1.seqmod; 838
  END seqmod; 839
840
  METHOD specify; 841
    RUN seqmod; 842
    RUN f1.specify; 843
    RUN f2.specify; 844
```

```

END specify; 845
846
METHOD reset; 847
  RUN clear; 848
  RUN specify; 849
END reset; 850
END flowsheet; 851
852
853
(* ***** *) 854
855
856
MODEL test_flowsheet REFINES flowsheet; 857
858
f1.stream.components := ['A', 'B', 'C', 'D']; 859
860
METHODS 861
  METHOD values; 862
    RUN reset; 863
    (* Initial configuration *) 864
    select_feed1 := TRUE; 865
    select_single1 := TRUE; 866
    select_cheapr1 := TRUE; 867
    select_single2 := TRUE; 868
869
    (* values of fixed variables *) 870
    f1.stream.state.Cpi['A'] := 0.04 {BTU/mole/K}; 871
    f1.stream.state.Cpi['B'] := 0.05 {BTU/mole/K}; 872
    f1.stream.state.Cpi['C'] := 0.06 {BTU/mole/K}; 873
    f1.stream.state.Cpi['D'] := 0.055 {BTU/mole/K}; 874
875
    col.heat_removed := 100 {BTU/s}; 876
    col.cost_factor := 0.7e-06 {dollar/kJ}; 877
878
    h1.heat_supplied := 200 {BTU/s}; 879
    h1.cost_factor := 8e-06 {dollar/kJ}; 880
881
    co2.heat_removed := 150 {BTU/s}; 882
    co2.cost_factor := 0.7e-06 {dollar/kJ}; 883
884
    f11.alpha['A'] := 12.0; 885
    f11.alpha['B'] := 10.0; 886
    f11.alpha['C'] := 1.0; 887
    f11.alpha['D'] := 6.0; 888
    f11.vap_to_feed_ratio := 0.9; 889
    f11.ave_alpha := 5.0; 890

```

```

                                                                    891
h2.heat_supplied := 180 {BTU/s};                                     892
h2.cost_factor   := 8e-06 {dollar/kJ};                             893
                                                                    894
sp1.split[1] := 0.05;                                             895
                                                                    896
h3.heat_supplied := 190 {BTU/s};                                     897
h3.cost_factor   := 8e-06 {dollar/kJ};                             898
                                                                    899
m1.To := 298 {K};                                                 900
                                                                    901
f1.cost_factor := 0.026 {dollar/kg_mole};                         902
f2.cost_factor := 0.033 {dollar/kg_mole};                         903
                                                                    904
c1.cost_factor := 8.33333e-06 {dollar/kJ};                       905
c1.pressure_rate := 2.5;                                          906
                                                                    907
s1.cost_factor_work := 8.33333e-06 {dollar/kJ};                 908
s1.cost_factor_heat := 0.7e-06 {dollar/kJ};                     909
s1.pressure_rate := 2.5;                                          910
                                                                    911
r1.stoich_coef['A'] := -1;                                        912
r1.stoich_coef['B'] := -1;                                        913
r1.stoich_coef['C'] := 1;                                        914
r1.stoich_coef['D'] := 0;                                        915
r1.low_turnover := 0.0069 {kg_mole/s};                          916
                                                                    917
r2.stoich_coef['A'] := -1;                                        918
r2.stoich_coef['B'] := -1;                                        919
r2.stoich_coef['C'] := 1;                                        920
r2.stoich_coef['D'] := 0;                                        921
r2.high_turnover := 0.00828 {kg_mole/s};                        922
                                                                    923
                                                                    924
c2.cost_factor := 8.33333e-06 {dollar/kJ};                       925
c2.pressure_rate := 1.5;                                          926
                                                                    927
s2.cost_factor_work := 8.33333e-06 {dollar/kJ};                 928
s2.cost_factor_heat := 0.7e-06 {dollar/kJ};                     929
s2.pressure_rate := 1.5;                                          930
END values;                                                       931
                                                                    932
METHOD configuration2;                                           933
  (* alternative configuration *)                                  934
  select_feed1 := FALSE;                                         935
  select_single1 := FALSE;                                       936
```



```
select_cheapr1 := FALSE;          937
select_single2 := FALSE;          938
END configuration2;                939
                                   940
METHOD configuration3;             941
  (* alternative configuration *)   942
  select_feed1 := FALSE;          943
  select_single1 := TRUE;         944
  select_cheapr1 := TRUE;         945
  select_single2 := FALSE;        946
END configuration3;                947
END test_flowsheet;               948
(* ***** *)                    949
```

# CHAPTER 18 A SIMPLE CHEMICAL ENGINEERING FLOWSHEETING EXAMPLE

In this example we shall examine a model for a simple chemical engineering process flowsheet. The code listed below exists in the file in the ASCEND examples subdirectory entitled *simple\_fs.asc*. Except for some formatting changes to make it more presentable here, it is exactly as it is in the library version. Thus you could run this example by loading this file and using it and its corresponding script *simple\_fs.s*.

## 18.1 THE PROBLEM DESCRIPTION

This model is of a simple chemical engineering flowsheet. Studying it will help to see how one constructs more complex models in ASCEND. Models for more complex objects are typically built out of previously defined types each of which may itself be built of previously defined parts, etc. A flowsheet could, for example, be built of units and streams. A distillation column could itself be built out of trays and interconnecting streams.

Lines 40 to 56 in the code below give a diagram of the flowsheet we would like to model. This flowsheet is to convert species B into species C. B undergoes the reaction.



The available feed contains 5 mole percent of species A, a light contaminant that acts as an inert in the reactor. We pass this feed into the reactor where only about 7% of B converts per pass. Species C is much less volatile than B which is itself somewhat less volatile than A. Relative volatilities are 12, 10 and 1 respectively for A, B and C. Species A will build up if we do not let it escape from the system. We propose to do this by bleeding off a small portion (say 1 to 2%) of the B we recover and recycle back to the reactor.

The flowsheet contains a mixer where we mix the recycle with the feed, a reactor, a flash unit, and a stream splitter where we split off and remove some of the recycled species B contaminated with species A

Our goal is to determine the impact of the bleed on the performance of this flowsheet. We would also like to see if we can run the flash unit to get us fairly pure C as a bottom product from it.

The first type definitions we need for our simple flowsheet are for the variables we would like to use in our model. The ones needed for this example are all in the file *atoms.a4l*. Thus we will need to load *atoms.a4l* before we load the file containing the code for this model.

The following is the code for this model. We shall intersperse comments on the code within it.

## 18.2 THE CODE

As the code is in our ASCEND models directory, it has header information that we require of all such files included as one large comment extending over several lines. Comments are in the form (\* comment \*).

To assure that appropriate library files are loaded first, ASCEND has the REQUIRE statement, such as appears on line 61:

```
REQUIRE atoms.a4l
```

This statement causes the system to load the file *atoms.a4l* before continuing with the loading of this file. *atoms.a4l* in turn has a require statement at its beginning to cause *system.a4l* to be loaded before it is.

```
(*****\
    simple_fs.asc
    by Arthur W. Westerberg
    Part of the Ascend Library
This file is part of the Ascend modeling library.
Copyright (C) 1994
The Ascend modeling library is free software; you can redistribute
it and/or modify it under the terms of the GNU General Public License as
published by the Free Software Foundation; either version 2 of the
License, or (at your option) any later version.
The Ascend Language Interpreter is distributed in hope that it will be
useful, but WITHOUT ANY WARRANTY; without even the implied warranty of
MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE. See the GNU
```

General Public License for more details. 18

You should have received a copy of the GNU General Public License along  
with the program; if not, write to the Free Software Foundation, Inc.,  
675 Mass Ave, Cambridge, MA 02139 USA. Check the file named COPYING. 20  
21  
22

Use of this module is demonstrated by the associated script file  
simple\_fs.s. 24  
25

\\*\*\*\*\*) 26

(\*\*\*\*\*\ 28

\$Date: 97/02/20 18:54:21 \$ 29

\$Revision: 1.5 \$ 30

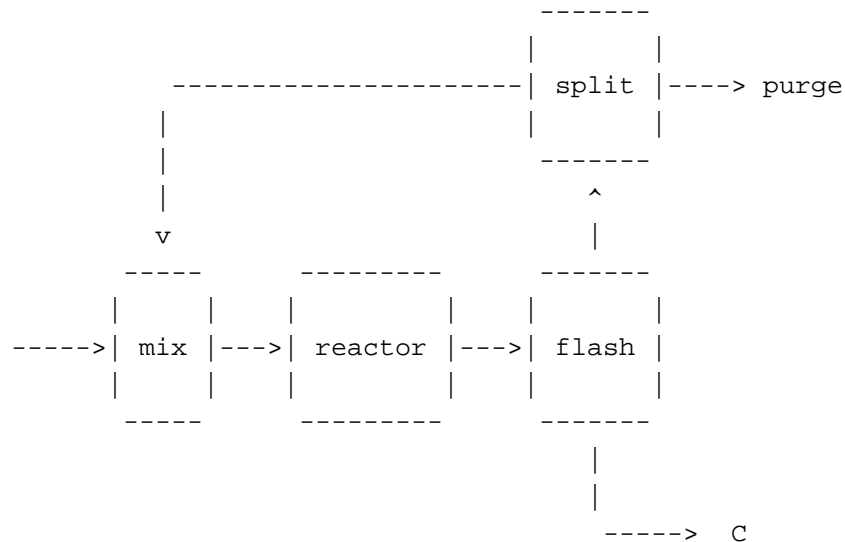
\$Author: mthomas \$ 31

\$Source: /afs/cs.cmu.edu/project/ascend/Repository/models/examples/  
simple\_fs.asc,v \$ 32

\\*\*\*\*\*) 33

(\* 34

The following example illustrates equation based modeling using the  
ASCEND system. The process is a simple recycle process. 36  
37  
38



This model requires: "system.a41"  
"atoms.a41" 57  
58

\*) 59

REQUIRE atoms.a41 60  
61

The first model we shall define is for defining a stream. In the document entitled [\\*\\*\\*\\*link to \\*\\*\\*\\* Equation-based Process Modeling\\*\\*\\*\\*](#) we argued the need to define a stream by maximizing the use of intensive variables and the equations interrelating them. Our problem here requires only the molar flows for the components as the problem definition provides us with all the physical properties as constants. Nowhere for this simple model do we seem to need temperatures, fugacities, etc. To maximize the use of intensive variables, we will use mole fractions and total molar flow to characterize a stream. We must include the equation that says the mole fractions add to unity. Our first model we call *mixture*.

```
( * ***** *) 62
MODEL mixture; 63
64
  components          IS_A set OF symbol_constant; 65
  y[components]      IS_A fraction; 66
67
  SUM[y[i] | i IN components] = 1.0; 68
69
METHODS 70
71
  METHOD clear; 72
    y[components].fixed := FALSE; 73
  END clear; 74
75
  METHOD specify; 76
    y[components].fixed := TRUE; 77
    y[CHOICE[components]].fixed := FALSE; 78
  END specify; 79
80
  METHOD reset; 81
    RUN clear; 82
    RUN specify; 83
  END reset; 84
85
END mixture; 86
87
```

Line 66 of the model for mixture defines a set of symbol constants. We will later include in this set one symbol constant giving a name for each of the species in the problem (A, B and C). Line 67 defines one mole fraction variable for each element in the set of components, while line 69 says these mole fractions must add to 1.0.

We add a methods section to our model to handle the flag setting which we shall need when making the problem well-posed -- i.e., as a problem having an equal number of unknowns as equations. We first have a method called `clear` which resets all the “fixed” flags for all the variables in this model to `FALSE`. This method puts the problem into a known state (all flags are `FALSE`). The second method is our selection of variables that we wish to fix if we were to solve the equations corresponding to a mixture model. There is only one equation among all the mole fraction variables so we set all but one of the flags to `TRUE`. The `CHOICE` function picks arbitrarily one of the members of the set *components*. For that element, we reset the fixed flag to `FALSE`, meaning that this one variable will be computed in terms of the values given to the others.

The reset method is useful as it runs first the `clear` method to put an instance of a mixture model into a known state with respect to its fixed flags, followed by running the `specify` method to set all but one of the fixed flags to `TRUE`.

These methods are not needed to create our model. To include them is a matter of modeling style, a style we consider to be good practice. The investment into writing these methods now has always been paid back in reducing the time we have needed to debug our final models.

The next model we write is for a stream, a model that will include a part we call *state* which is an instance of the type *mixture*.

```
( * ***** *) 88
MODEL molar_stream; 89
  components      IS_A set OF symbol_constant; 90
  state           IS_A mixture; 91
  Ftot,f[components]IS_A molar_rate; 92
  components, state.componentsARE_THE_SAME; 93
  FOR i IN components CREATE 94
    f_def[i]: f[i] = Ftot*state.y[i]; 95
  END; 96
METHODS 97
  METHOD clear; 98
    RUN state.clear; 99
    Ftot.fixed := FALSE; 100
  101
  102
  103
  104
  105
  106
```

```

        f[components].fixed:= FALSE;                               107
    END clear;                                                    108
                                                                    109
    METHOD seqmod;                                                110
        RUN state.specify;                                       111
        state.y[components].fixed:= FALSE;                       112
    END seqmod;                                                  113
                                                                    114
    METHOD specify;                                              115
        RUN seqmod;                                             116
        f[components].fixed:= TRUE;                              117
    END specify;                                                118
                                                                    119
    METHOD reset;                                               120
        RUN clear;                                             121
        RUN specify;                                           122
    END reset;                                                  123
                                                                    124
    METHOD scale;                                               125
        FOR i IN components DO                                  126
            f[i].nominal := f[i] + 0.1{mol/s};                 127
        END;                                                    128
        Ftot.nominal := Ftot + 0.1{mol/s};                     129
    END scale;                                                  130
                                                                    131
END molar_stream;                                             132
                                                                    133

```

We define our stream over a set of components. We next include a part which is of type mixture and call it *state* as mentioned above. We also include a variable entitled *Ftot* which will represent the total molar flowrate for the stream. For convenience -- as they are not needed, we also include the molar flows for each of the species in the stream. We realize that the components defined within the part called *state* and the set of components we just defined for the stream should be the same set. We force the two sets to be the same set with the `ARE_THE_SAME` operator.

We next write the equations that define the individual molar flows for the components in terms of their corresponding mole fractions and the total flowrate for the stream. Note, the equations that says the mole fractions add to unity in the definition of the state forces the total of the individual flowrates to equal the total flowrate. Thus we do not need to include an equation that says the molar flowrates for the species add up to the total molar flowrate for the stream.

We again write the methods we need for handling flag setting. We leave it to the reader to establish that the specify method produces a well-posed instance involving the same number of variables to be computed as equations available to compute them. The scale method is there as we may occasionally wish to rescale the nominal values for our flows to reflect the values we are computing for them. Poor scaling of variables can lead to numerical difficulties for really large models. This method is there to reduce the chance we will have poor scaling.

Note that the nominal values for the remaining variables -- the mole fractions -- are unity. It does not need to be recomputed as unity is almost always a good nominal value for each of them.

Our next model is for the first of our unit operations. Each of these will be built of streams and equations that characterize their behavior. The first models a mixer. It can have any number of feed streams, each of which is a molar stream. We require the component set for each of the feed streams and the output stream from the unit to be the same set. Finally we write a component material balance for each of the species in the problem, where we sum the flows in each of the feeds to give the flow in the output stream, *out*.

```
( * ***** *) 134
                                                                135
MODEL mixer; 136
                                                                137
    n_inputs          IS_A  integer_constant; 138
    feed[1..n_inputs], out  IS_A  molar_stream; 139
                                                                140
    feed[1..n_inputs].components, 141
    out.components          ARE_THE_SAME; 142
                                                                143
    FOR i IN out.components CREATE 144
        cmb[i]: out.f[i] = SUM[feed[1..n_inputs].f[i]]; 145
    END; 146
                                                                147
METHODS 148
                                                                149
    METHOD clear; 150
        RUN feed[1..n_inputs].clear; 151
        RUN out.clear; 152
    END clear; 153
                                                                154
    METHOD seqmod; 155
    END seqmod; 156
                                                                157
```



```

METHOD specify;                                158
  RUN seqmod;                                  159
  RUN feed[1..n_inputs].specify;              160
END specify;                                    161
                                                162
METHOD reset;                                   163
  RUN clear;                                   164
  RUN specify;                                 165
END reset;                                       166
                                                167
METHOD scale;                                   168
  RUN feed[1..n_inputs].scale;                 169
  RUN out.scale;                               170
END scale;                                       171
                                                172
END mixer;                                       173

```

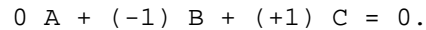
The *METHOD clear* sets all the fixed flags for the parts of this model to false by running each of their clear methods (i.e., for all the feeds and for the stream out). If this model had introduced any new variables, their fixed flags would have been set to FALSE here.

We will implement the method to make the model well posed into two parts: *seqmod* (stands for “sequential modular” which is the mindset we use to get a unit well-posed) and *specify*. The first we shall use within any unit operation to fix exactly enough fixed flags for a unit such that, if we also make the feed streams to it well-posed, the unit will be well-posed. For a mixer unit, the output stream results simply from mixing the input streams; there are no other variables to set other than those for the feeds. Thus the *seqmod* method is empty. It is here for consistency with the other unit operation models we write next. The *METHOD specify* makes this model well-posed by calling the *seqmod* method and then the *specify* method for each of the feed streams. No other flags need be set to make the model well-posed.

*METHOD reset* simply runs *clear* followed by *specify*. Running this sequence of method will make the problem well-posed no matter which of the fixed flags for it are set to TRUE before running *reset*. Finally, flowrates can take virtually any value so we can include a *scale* method to scale the flows based on their current values.

The next model is for a very simple ‘degree of conversion’ reactor. The model defines a turnover rate which is the rate at which the reaction as written proceeds (e.g., in moles/s). For example, here our reaction will be  $B \rightarrow C$ . A turnover rate of 3.7 moles/s would mean that 3.7 moles/s of B would convert to 3.7 moles/s of C. The vector *stoich\_coef* has one

entry per component. Here there will be three components when we test this model so the coefficients would be 0, -1, 1 for the reaction



Reactants have a negative coefficient, reactants a positive one. The material balance to compute the flow out for each of the components sums the amount coming in plus that created by the reaction.

```
( * ***** *) 174
MODEL reactor; 175
176
177
    feed, out          IS_A molar_stream; 178
    feed.components, out.components ARE_THE_SAME; 179
180
    turnover IS_A molar_rate; 181
    stoich_coef[feed.components] IS_A factor; 182
183
    FOR i IN feed.components CREATE 184
        out.f[i] = feed.f[i] + stoich_coef[i]*turnover; 185
    END; 186
187
METHODS 188
189
    METHOD clear; 190
        RUN feed.clear; 191
        RUN out.clear; 192
        turnover.fixed := FALSE; 193
        stoich_coef[feed.components].fixed := FALSE; 194
    END clear; 195
196
    METHOD seqmod; 197
        turnover.fixed := TRUE; 198
        stoich_coef[feed.components].fixed := TRUE; 199
    END seqmod; 200
201
    METHOD specify; 202
        RUN seqmod; 203
        RUN feed.specify; 204
    END specify; 205
206
    METHOD reset; 207
        RUN clear; 208
        RUN specify; 209
    END reset; 210
```

```

METHOD scale; 211
  RUN feed.scale; 212
  RUN out.scale; 213
  turnover.nominal := turnover.nominal+0.0001 {kg_mole/s}; 215
END scale; 216
217
END reactor; 218
219

```

The *METHOD clear* first directs all the parts of the reactor to run their *clear* methods. Then it sets the fixed flags for all variables introduced in this model to FALSE.

Assume the feed to be known. We introduced one stoichiometric coefficient for each component and a turnover rate. To make the output stream well-posed, we would need to compute the flows for each of the component flows leaving. That suggests the material balances we wrote are all needed to compute these flows. We would, therefore, need to set one fix flag to TRUE for each of the variables we introduced, which is what we do in the *METHOD seqmod*. Now when we run *seqmod* and then the *specify* method for the feed, we will have made this model well-posed, which is what we do in the *METHOD specify*.

The flash model that follows is a constant relative volatility model. Try reasoning why the methods attached are as they are.

```

( * ***** *) 220
MODEL flash; 221
222
  feed,vap,liqIS_Amolar_stream; 223
224
  feed.components, 225
  vap.components, 226
  liq.components ARE_THE_SAME; 227
228
  alpha[feed.components], 229
  ave_alpha IS_A factor; 230
231
  vap_to_feed_ratio IS_A fraction; 232
233
  vap_to_feed_ratio*feed.Ftot = vap.Ftot; 234
235
FOR i IN feed.components CREATE 236
  cmb[i]: feed.f[i] = vap.f[i] + liq.f[i]; 237
238

```

```

    eq[i]:  vap.state.y[i]*ave_alpha = alpha[i]*liq.state.y[i];      239
  END;                                                                240
                                                                    241
METHODS                                                                242
                                                                    243
METHOD clear;                                                         244
  RUN feed.clear;                                                    245
  RUN vap.clear;                                                     246
  RUN liq.clear;                                                     247
  alpha[feed.components].fixed   := FALSE;                          248
  ave_alpha.fixed                 := FALSE;                          249
  vap_to_feed_ratio.fixed         := FALSE;                          250
END clear;                                                            251
                                                                    252
METHOD seqmod;                                                       253
  alpha[feed.components].fixed   := TRUE;                            254
  vap_to_feed_ratio.fixed         := TRUE;                            255
END seqmod;                                                           256
                                                                    257
METHOD specify;                                                      258
  RUN seqmod;                                                         259
  RUN feed.specify;                                                  260
END specify;                                                          261
                                                                    262
METHOD reset;                                                         263
  RUN clear;                                                          264
  RUN specify;                                                        265
END reset;                                                            266
                                                                    267
METHOD scale;                                                         268
  RUN feed.scale;                                                    269
  RUN vap.scale;                                                     270
  RUN liq.scale;                                                     271
END scale;                                                            272
                                                                    273
END flash;                                                            274
                                                                    275
( * ***** * )                                                    276
                                                                    277

```

The final unit operation model is the splitter. The trick here is to make all the states for all the output streams the same as that of the feed. This move makes the compositions all the same and introduces only one equation to add those mole fractions to unity. The rest of the model should be evident.

```

MODEL splitter; 278
                279
    n_outputs    IS_A integer_constant; 280
    feed, out[1..n_outputs ] IS_A molar_stream; 281
    split[1..n_outputs] IS_A fraction; 282
                283
    feed.components, out[1..n_outputs].components ARE_THE_SAME; 284
                285
    feed.state, 286
    out[1..n_outputs].state ARE_THE_SAME; 287
                288
    FOR j IN [1..n_outputs] CREATE 289
        out[j].Ftot = split[j]*feed.Ftot; 290
    END; 291
                292
    SUM[split[1..n_outputs]] = 1.0; 293
                294
METHODS 295
                296
    METHOD clear; 297
        RUN feed.clear; 298
        RUN out[1..n_outputs].clear; 299
        split[1..n_outputs-1].fixed:=FALSE; 300
    END clear; 301
                302
    METHOD seqmod; 303
        split[1..n_outputs-1].fixed:=TRUE; 304
    END seqmod; 305
                306
    METHOD specify; 307
        RUN seqmod; 308
        RUN feed.specify; 309
    END specify; 310
                311
    METHOD reset; 312
        RUN clear; 313
        RUN specify; 314
    END reset; 315
                316
    METHOD scale; 317
        RUN feed.scale; 318
        RUN out[1..n_outputs].scale; 319
    END scale; 320
                321
END splitter; 322
                323

```

```
( * ***** *) 324
325
```

Now we shall see the value of writing all those methods for our unit operations (and for the models that we used in creating them). We construct our flowsheet by saying it includes a mixer, a reactor, a flash unit and a splitter. The mixer will have two inputs and the splitter two outputs. The next few statements configure our flowsheet by making, for example, the output stream from the mixer and the feed stream to the reactor be the same stream.

The methods are as simple as they look. This model does not introduce any variables nor any equations that are not introduced by its parts. We simply ask the parts to clear their variable fixed flags.

To make the flowsheet well-posed, we ask each unit to set sufficient fixed flags to TRUE to make itself well posed were its feed stream well-posed (now you can see why we wanted to create the methods *seqmod* for each of the unit types.) Then the only streams we need to make well-posed are the feeds to the flowsheet, of which there is only one. The remaining streams come out of a unit which we can think of computing the flows for it.

```
MODEL flowsheet; 326
327
  m1          IS_A  mixer; 328
  r1          IS_A  reactor; 329
  fl1        IS_A  flash; 330
  spl        IS_A  splitter; 331
332
(* define sets *) 333
334
  m1.n_inputs  ==2; 335
  spl.n_outputs ==2; 336
337
(* wire up flowsheet *) 338
339
  m1.out, r1.feed  ARE_THE_SAME; 340
  r1.out, fl1.feed ARE_THE_SAME; 341
  fl1.vap, spl.feed ARE_THE_SAME; 342
  spl.out[2], m1.feed[2] ARE_THE_SAME; 343
344
METHODS 345
346
  METHOD clear; 347
    RUN m1.clear; 348
```

```

        RUN r1.clear;                                349
        RUN f11.clear;                               350
        RUN sp1.clear;                               351
    END clear;                                       352
                                                    353
    METHOD seqmod;                                    354
        RUN m1.seqmod;                               355
        RUN r1.seqmod;                               356
        RUN f11.seqmod;                             357
        RUN sp1.seqmod;                             358
    END seqmod;                                       359
                                                    360
    METHOD specify;                                   361
        RUN seqmod;                                  362
        RUN m1.feed[1].specify;                     363
    END specify;                                       364
                                                    365
    METHOD reset;                                     366
        RUN clear;                                   367
        RUN specify;                                 368
    END reset;                                       369
                                                    370
    METHOD scale;                                     371
        RUN m1.scale;                                372
        RUN r1.scale;                                373
        RUN f11.scale;                               374
        RUN sp1.scale;                               375
    END scale;                                       376
                                                    377
END flowsheet;                                       378
                                                    379
(* ***** *)                                       380
                                                    381

```

We have created a flowsheet model above. If you look at the reactor model, we require the use specify the turnover rate for the reaction. We may have no idea of a suitable turnover rate. What we may have an idea about is the conversion of species B in the reactor; for example, we may know that about 7% of the B entering the reactor may convert. How can we alter our model to allow for us to say this about the reactor and not be required to specify the turnover rate? In a sequential modular flowsheeting system, we would use a “computational controller.” We shall create a model here that gives us this same functionality. Thus we call it a “controller.” There are many ways to construct this model. We choose here to create a model here that has a flowsheet as a part of it. We introduce a variable conv which will

indicate the fraction conversion of any one of the components which we call the `key_component` here. For that component, we add a material balance based on the fraction of it that will convert. We added one new variable and one new equation so, if the flowsheet is well-posed, so will our controller be well-posed. However, we want to specify the conversion rather than the turnover rate. The *specify* method first asks the flowsheet `fs` to make itself well-posed. Then it makes this one trade: fixing `conv` and releasing the turnover rate.

```

MODEL controller;                                     382
                                                    383
    fs                IS_A  flowsheet;               384
    conv              IS_A  fraction;                 385
    key_components    IS_A  symbol_constant;          386
    fs.r1.out.f[key_components] = (1 - conv)*fs.r1.feed.f[key_components]; 387
                                                    388
METHODS                                               389
                                                    390

METHOD clear;                                         391
    RUN fs.clear;                                     392
    conv.fixed:=FALSE;                               393
END clear;                                           394
                                                    395

METHOD specify;                                       396
    RUN fs.specify;                                   397
    fs.r1.turnover.fixed:=FALSE;                     398
    conv.fixed:=TRUE;                                 399
END specify;                                         400
                                                    401

METHOD reset;                                        402
    RUN clear;                                       403
    RUN specify;                                     404
END reset;                                           405
                                                    406

METHOD scale;                                        407
    RUN fs.scale;                                    408
END scale;                                           409
                                                    410

END controller;                                     411
                                                    412
(* ***** *)                                       413
                                                    414

```

We now would like to test our models to see if they work. How can we write test for them? We can create test models as we do below.



To test the flowsheet model, we create a `test_flowsheet` model that refines our previously defined flowsheet model. To refine the previous model, means this model includes all the statements made to define the flowsheet model plus those statements that we now provide here. So this model is a flowsheet but with its components specified to be 'A', 'B', and 'C'. We add a new method called *values* in which we specify values for all the variables we intend to fix when we solve. We can also provide values for other variables; these will be used as the initial values for them when we start to solve. We see all the variables being given values with the units specified. The units must be specified in ASCEND. ASCEND will interpret the lack of units to mean the variable is unitless. If it is not, then you will get a diagnostic from ASCEND telling you that you have written a dimensionally inconsistent relationship.

Note we specify the molar flows for the three species in the feed. Given these flows, the equations for the stream will compute the total flow and then the mole fractions for it. Thus the feed stream is fully specified with these flows.

We look at the `seqmod` method for each of the units to see the variables to which we need to give values here.

```

MODEL test_flowsheet REFINES flowsheet;                                415
                                                                           416
    m1.out.components:==['A','B','C'];                                  417
                                                                           418
METHODS                                                                    419
                                                                           420
METHOD values;                                                            421
    m1.feed[1].f['A']           := 0.005 {kg_mole/s};                    422
    m1.feed[1].f['B']           := 0.095 {kg_mole/s};                    423
    m1.feed[1].f['C']           := 0.0   {kg_mole/s};                    424
                                                                           425
    r1.stoich_coef['A']         := 0;                                     426
    r1.stoich_coef['B']         := -1;                                    427
    r1.stoich_coef['C']         := 1;                                     428
    r1.turnover                  := 3 {kg_mole/s};                       429
                                                                           430
    f11.alpha['A']              := 12.0;                                  431
    f11.alpha['B']              := 10.0;                                  432
    f11.alpha['C']              := 1.0;                                  433
    f11.vap_to_feed_ratio       := 0.9;                                  434
    f11.ave_alpha                := 5.0;                                  435
                                                                           436
    sp1.split[1]                 := 0.01;                                 437

```

```

f11.liq.Ftot:=m1.feed[1].f['B'];
END values;
END test_flowsheet;
(* ***** *)

```

Finally we would like to test our controller model. Again we write our test model as a refinement of the model to be tested. The test model is, therefore, a controller itself. We make our fs model inside our test model into a test\_flowsheet, making it a more refined type of part than it was in the controller model. We can do this because the test\_controller model is a refinement of the flowsheet model which fs was previously. A test\_flowsheet is, as we said above, a flowsheet. We create a values method by first running that we wrote for the test\_flowsheet model and then add a specification for the conversion of B in the reactor.

```

MODEL test_controller REFINES controller;
fs      IS_REFINED_TOtest_flowsheet;
key_components := 'B';
METHODS
METHOD values;
  RUN fs.values;
  conv      := 0.07;
END values;
END test_controller;
(* ***** *)

```



# CHAPTER 19 THE ASCEND PREDEFINED COLLECTION OF MODELS

The ASCEND system has two subdirectories containing models we and others have previously defined. We have labeled the first subdirectory **libraries** and the second **examples**. In the **libraries** subdirectory are several different files, each containing a number of useful type definitions which we can use to construct larger models in ASCEND. The **examples** subdirectory contains a number of complete ASCEND models ready for us to execute. One can examine and execute these examples when learning how to model in ASCEND.

## **system.a41**

The file called *system.a41* in the libraries subdirectory must always be loaded first in the ASCEND system. It is automatically loaded when one starts the ASCEND system. However, the *delete all types* command will delete all type definitions including the ones in this file. If you have deleted all types, always reload this file first using the *Read* instruction in the Library tool set.

## **atoms.a41**

The simplest collection of previously defined types are those which define the kinds of constants, parameters and variables we are likely to use in constructing an engineering or scientific model. A file called *atoms.a41* located in the libraries subdirectory has over 125 types of constants, parameters and variables. Following are three of the definitions it contains.

```

CONSTANT critical_compressibility REFINES
    real_constant DIMENSIONLESS;                                1
                                                                2
UNIVERSAL CONSTANT speed_of_light                             3
REFINES real_constant ::= 1{LIGHT_C};                          4
                                                                5
ATOM volume REFINES solver_var                                 6
    DIMENSION L^3                                              7
    DEFAULT 100.0{ft^3};                                       8
    lower_bound := 0.0{ft^3};                                  9
    upper_bound := 1e50{ft^3};                                10
    nominal := 100.0{ft^3};                                    11
END volume;                                                    12

```

Note that the first and third include a statement of the dimensionality of the item being defined. For example critical compressibility is dimensionless while the dimensions for volume are  $L^3$  (i.e., length cubed). The ASCEND system supports nine basic dimensions as defined for the standards defining the SI system of units. Dimensions differ from units in that *length* is a dimension while *feet* is a set of units one may use to express a length. Dimensions in ASCEND are L (length: typical units being ft, m), M (mass: kg, lbm), T (time: s, yr), E (electric current: amp), Q (quantity: mole), TMP (temperature: K, R), LUM (luminous intensity: candela), P (plane angle: radian) and S (solid angle: steradian). We have also included the tenth dimension C (currency: USdollar) so one can express cost. If you wish to express cost in a variety of different currencies (e.g., USdollars, UKpounds), you will have to define the conversion rates.

(See the manual entitled (\*\*\*\*[hyperlink to the document entitled \\*\\*\\*\\*](#) The ASCEND Language Syntax and Semantics for more information on dimensionality and units.)

#### **Typical use of library files**

One will typically create models in the ASCEND system by including one or more of the library files available. Almost certainly the file *atoms.a4l* will become a part of any engineering or scientific model.

It would be useful for you to view this and a few of the other library files using a text editor such as xemacs to see what libraries we do have available.

#### **Examples and scripts**

The examples subdirectory in ASCEND has a number of complete ASCEND models. Each model is in two parts: the .a4c file containing the model definition and the .a4s file containing a script which one can use to execute the model. An example is the model *simple\_fs.a4c* along with its script *simple\_fs.a4s*.

Each of the example files indicates which of the library files one must load and the order in which to load them before loading the example file. If you fail to load a library file, you will experience a large number of diagnostic messages indicating there are missing type definitions.

# CHAPTER 20 THE ASCEND IV LANGUAGE

## SYNTAX AND SEMANTICS

Benjamin Allan<sup>1</sup>

Arthur W. Westerberg<sup>1</sup>

Department of Chemical Engineering  
and the Engineering Design Research Center /  
Institute for Complex Engineered Systems

Carnegie Mellon University

We shall present an informal description of the ASCEND IV language. Being informal, we shall usually include examples and descriptions of the intended semantics along with the syntax of the items. At times the inclusion of semantics will seem to anticipate later definitions. We do this because we would also like this chapter to be used as a reference for the ASCEND language even after one generally understands it. Often one will need to clarify a point about a particular item and will not wish to have to search in several places to do so.

*Syntax* is the form or structure for the statements in ASCEND, where one worries about the exact words one uses, their ordering, the punctuation, etc. *Semantics* describe the meaning of a statement.

To distinguish between syntax and semantics, consider the statement

```
y IS_A fraction;
```

Rules on the syntax for this statement tell us we need a user supplied instance name, *y*, followed by the ASCEND operator *IS\_A*, followed by a type name (*fraction*). The statement terminates with a semicolon. The statement semantics says we are declaring the existence of an instance,

---

1. The ASCEND language has evolved from the combined efforts of several generations of users and implementors. We wish to particularly acknowledge the contributions of ASCEND III implementors Kirk Abbott, Tom Epperly, Peter Piela, Boyd Safrit, Karl Westerberg, and Joe Zaher, and of the ASCEND IV crew: Vicente Rico-Ramirez, Mark Thomas and Ken Tyner.

locally named  $y$ , of the type fraction as a part within the current model definition and it is to be constructed when an instance of the current model definition is constructed.

The syntax for a computer language is often defined by using a Bachus-Naur formal (BNF) description. The complete YACC and FLEX description of the language described (as presently implemented) is available by FTP<sup>2</sup> and via the World Wide Web<sup>3</sup>. The semantics of a very high level modeling language such as ASCEND IV are generally much more restrictive than the syntax. For this reason we do not include a BNF description in this paper. ASCEND IV is an experiment. The language is under constant scrutiny and improvement, so this document is under constant revision. Contact the authors for the latest version.

## 20.1 PRELIMINARIES

We will start off with some background information and some tips that make the rest of the chapter easier to read. ASCEND is an object-oriented (OO) language for hierarchical modeling that has been somewhat specialized for mathematical models. Most of the specialization is in the implementation and the user interface rather than the language definition.

We feel the single most distinguishing feature of mathematical models is that solving them efficiently requires that the solving algorithms be able to address the entire problem either simultaneously or in a decomposition of the natural problem structure that the algorithm determines is best for the machine(s) in use. In the ASCEND language object-orientation is used to organize natural structures and make them easier to understand. It is not used to hide the details of the objects. The user (or machine) is free to ignore uninteresting details, and the ASCEND environment provides tools for the runtime suppression of these.

ASCEND is beginning its 4th generation. Some features we will describe are not yet implemented (some merely speculative) and these are clearly marked (\* 4+ \*). Any feature not marked (\* 4+ \*) has been completely implemented, and thus any mismatch between the description given here and the software we distribute is a bug we want you to tell us about.

The syntax and semantics of ASCEND may seem at first a bit unusual. However, do not be afraid to just try what comes naturally if what we write here is unclear. The parser and compiler of ASCEND IV really will help you get things right. Of course if what

---

2. In the directory [ftp.cs.cmu.edu:project/ascend/gnu-ascend/](ftp://ftp.cs.cmu.edu/project/ascend/gnu-ascend/) see the file README.  
3. <http://www.cs.cmu.edu/afs/cs.cmu.edu/project/ascend/ftp/gnu-ascend/README>

we write here is unclear, please ask us about it because we aim to continuously improve both this document and the language system it describes.

We will describe, starting in Section 20.1.2, the higher level concepts of ASCEND, but first some important punctuation rules.

ASCEND is cAsE sensitive!

The keywords that are shown capitalized (or in lower case) in this chapter are that way because ASCEND is case sensitive. IS\_A is an ASCEND keyword; isa, Is\_a, and all the other permutations you can think of are NOT equivalent to IS\_A. In declaring new types of models and variables the user is free to use any style of capitalization he or she may prefer, however, they must remain consistent or undefined types and instances will result.

This case restriction makes our code very readable, but hard to type without a smart editor. We have kept the case-sensitivity because, like all mathematicians, we find ourselves running out of good variable names if we are restricted to a 26 letter alphabet. We have developed smart add-ins for two UNIX editors, EMACS and vi, for handling the upper case keywords and some other syntax elements. The use of these editors is described in another chapter.

The ASCEND IV parser is very picky and pedantic. It also tries to give helpful messages and occasionally even suggestions. New users should just dive in and make errors, letting the system help them learn how to avoid errors.

### 20.1.1 PUNCTUATION

This section covers both the punctuation that must be understood to read this document and the punctuation of ASCEND code.

**keywords:**

ASCEND keywords and type names are given in the left column in **bold** format. It is generally clear from the main text which are keywords and which are type names.

Minor items:

Minor headings that are helpful in finding details are given in the left column in underline format.

Tips:

Special notes and hints are sometimes placed on the left.

\*3\*:

This indicates that what follows is specific to ASCEND IIIc and may disappear in a future version of ASCEND IV. Generally ASCEND IV will provide some equivalent functionality at 1/10th of the ASCEND III price.



- \*4\* This indicates that what follows is specific to ASCEND IV and may not be available in ASCEND IIIc. Generally ASCEND III may provide some very klugey equivalent functionality, often at a very high price in terms of increased compilation time or debugging difficulty.
- \*4+\* ASCEND IV functionality that is not fully implemented at the time of this writing. The precise syntax of the final implementation may vary slightly from what is presented here. A revision of this document will be made at the time of implementation.
- LHS: Left Hand Side. Abbreviation used frequently.
- RHS: Right Hand Side. Abbreviation used frequently.
- Simple Names: In ASCEND simple names are made of the characters a through z, A through Z, \_, (\*4+\*: \$). The underscore is used as a letter, but it cannot be the first letter in a name. The "\$" character is used exclusively as the first character in the name of system defined built-in parts. "\$" is explained in more detail in Section 20.6.2. Simple names should be no more than 80 characters long.
- Compound names: Compound names are simple names strung together with dots (.). See the description of "." below.
- Groupings:
- « » In documentation optional fields are surrounded by these markers.
- ( \* \*) Comment. \*3\* Anything inside these is a comment. Comments DO NOT nest in ASCEND IIIc. Comments may extend over many lines. \*4\* Comments DO nest in ASCEND IV.
- ( ) Rounded parentheses. Used to enclose arguments for functions or models where the order of the arguments matters. Also used to group terms in complex arithmetic, logical, or set expressions where the order of operations needs to be specified.
- Efficiency tip: The compiler can simplify relation definitions in a particularly efficient manner if constants are grouped together.
- { } Curly braces. Used to enclose units. For example, 1 {kg\_mole/s}. Also used to enclose the body of annotations. **Note:** Curly braces are also used in TCL, the language of the ASCEND user interface, about which we will say more in another chapter.

- [ ] Square brackets. Used to enclose sets or elements of sets. Examples: `my_integer_set := [1,2,3]`, demonstrates the use of square brackets in the assignment of a set. `My_array[1]` demonstrates the use of square brackets in naming an array object indexed over an integer set which includes the element 1.
- .
- Dot. The dot is used, as in PASCAL and C, to construct the names of nested objects. Examples: if object a has a part b, then the way to refer to b is as a.b. `Tray[1].vle` shows a dot following a square bracket; here `Tray[1]` has a part named vle.
- .. Dot-dot or double dot. Integer range shorthand. For example, `my_integer_set := [1,2,3]` and `my_integer_set := [1..3]` are equivalent. If .. appears in a context requiring (), such as the ALIASES/IS\_A statement, then the range is expanded and ordered as we would naturally expect.
- :
- Colon. A separator used in various ways, principally to set the name of an arithmetic relation apart from the definition.
- ::
- Double colon. A separator used in the methods section for accessing methods defined on types other than the type the method is part of. Explained in Section 20.4.
- ;
- Semicolon. The separator of statements.

## 20.1.2 BASIC ELEMENTS

Boolean value TRUE or FALSE. Can't get much simpler, eh? In the language definition TRUE and FALSE do not map to 1 and 0 or any other type of numeric value. (In the implementation, of course, they do.)

User interface tip: The ASCEND user interface programmers have found it very convenient, however, to allow T/F, 1/0, Y/N, and other obvious boolean conventions as interactive input when assigning boolean values. We are lazy users.

Integer value A signed whole number up to the maximum that can be represented by the computer on which one is running ASCEND. MAX\_INTEGER is machine dependent. Examples are:

123  
-5

Typically, 2147483647. MAX\_INTEGER

Real value

ASCEND represents reals almost exactly as any other mathematically oriented programming language does. The mantissa has an optional negative sign followed by a string of digits and at most one decimal point. The exponent is the letter *e* followed by an integer. The number must not exceed the largest the computer is able to handle. There can be no blank characters in a real. `MAX_REAL` is machine dependent. The following are legitimate reals in ASCEND:

```
-1
1.2
1.3e-2
7.888888e+34
.6E21
```

Normally `MAX_REAL` is about 1.79E+308.

`MAX_REAL`

while the following are not:

```
1. 2 (*contains a blank within it*)
1.3e2.0 (*exponent has a decimal in it*)
+1.3 (*contains illegal unary + sign*)
```

Reals stored in SI units

We store all real values as double precision numbers in the MKS system of units. This eliminates many common errors in the modeling of physical systems. Since we also place the burden of scaling equations on system routines and a simple modeling methodology, the internal units are not of concern to most users.

Dimensionality:

Real values have dimensionality such as length/time for velocity. Dimensionality is to be distinguished from the units such as ft/s. ASCEND takes care of mapping between units and dimensions. A value without units (this includes integer values) is taken to be dimensionless. Dimensionality is built up from the following base dimensions:

Name	<u>definition</u> typical units
<b>L</b>	lengthmeter, m
<b>M</b>	masskilogram, kg
<b>T</b>	timesecond, s
<b>E</b>	electric currentampere, A

<b>Q</b>	quantitymole, mole
<b>TMP</b>	temperatureKelvin, K
<b>LUM</b>	luminous intensitycandela, cd
<b>P</b>	plane anglesteradian, rad
<b>S</b>	solid anglesteradian, sr
<b>C</b>	currencycurrency, CR

The atom and constant definitions in the library illustrate the use of dimensionality.

Dimensions may be any combination of these symbols along with rounded parentheses, (), and the operators \*, ^ and /. Examples include  $M/T$  or  $M*L^2/T^2/TMP$  {this latter means  $(M*(L^2)/(T^2))/TMP$ }. The second operand for the “to the power” operator, ^, must be an integer value (e.g., -2 or 3) because fractional powers are physically undefined.

If the dimensionality for a real value is undefined, then ASCEND gives it a wild card dimensionality. If ASCEND can later deduce its dimensionality from its use in a model definition it will do so. For example consider the real variable  $a$ , suppose  $a$  has wild card dimensionality,  $b$  has dimensionality of  $L/T$ . Then the statement:

Example of a dimensionally consistent equation.

$$a + b = 3 \text{ \{ft/s\}};$$

requires that  $a$  have the same dimensionality as the other two terms, namely,  $L/T$ . ASCEND will assign this dimensionality to  $a$ . The user will be warned of dimensionally inconsistent equations.

### Unit expression

A unit expression may be composed of any combination of unit names defined by the system and any numerical constants combined with times (\*), divide(/) and “to the power” (^) operators. The RHS of ^ must be an integer. Parentheses can be used to group subexpressions EXCEPT a divide operator may not be followed by a grouped subexpression.

So, {kg/m/s} is fine, but {kg/(m\*s)} is not. Although the two expressions are mathematically equivalent, it makes the system programming and output formatting easier to code and faster to execute if we disallow expressions of the latter sort.

The units understood by the system are defined in Chapter 21. Note that several “units” defined are really values of interesting constants in SI, e.g.  $R ::= 1\{\text{GAS\_C}\}$  yields the correct value of the thermodynamic gas constant. Users can define additional units.

## Units

A unit expression must be enclosed in curly braces `{}`. When a real number is used in a mathematical expression in ASCEND, it must have a set of units expressed with it. If it does not, ASCEND assumes the number is dimensionless, which may not be the intent of the modeler. An example is shown in the dimensionally consistent equation above where the number 3 has the units `{ft/s}` associated with it.

Examples:

```
{kg_mole/s/m} same as {(kg_mole/s)/m}
{m^3/yr}
{3/100*ft} same as {0.03*ft}
{s^-1} same as {1/s}
```

Illegal unit examples are

```
{m/(K*kg_mole)} grouped subexpression used in denominator (should
be written {m/K/kg_mole})
{m^3.5} power must be integer.
```

## Symbol Value

The format for a symbol is that of an arbitrary character string enclosed between two single quotes. There is no way to embed a single quote in a symbol: we are not in the escape sequence business at this time. The following are legal symbols in ASCEND:

```
'H2O'
'r1'
'bill said,"foo" to who?'
```

while the following are not legal symbol values:

```
"ethanol" (double quotes not allowed)
water (no single quotes given)
'i can't do this' (no embedded quotes)
```

There is an arbitrary upper limit to the number of characters in a symbol (something like 10,000) so that we may detect a missing close quote in a bad input file without crashing.

Sets values

Sets values are lists of elements, all of type `integer_constant` or all of type `symbol_constant`, enclosed between square brackets []. The following are examples of sets:

```
['methane', 'ethane', 'propane']
[1..5, 7, 15]
[2..n_stages]
[1, 4, 2, 1, 16]
[]
```

We will say more about sets in 20.2.2.

The value range `1..5` is an allowable shorthand for the integers 1, 2, 3, 4 and 5 while the value range `2..n_stages` (where `n_stages` must be of type `integer`) means all integers from 2 to `n_stages`. If `n_stages` is less than 2, then the third set is empty. The repeated occurrence of 1 in the fourth set is ignored. The fifth set is the empty set.

We use the term *set* in an almost pure mathematical sense. The elements have no order. One can only ask two things of a set: (1) if an element is a member of it and (2) its cardinality (`CARD(set)`). Repeated elements used in defining a set are ignored. The elements of sets **cannot** themselves be sets in ASCEND; i.e., there can be no sets of set.

Sets are unordered.

A set of integers may appear to be ordered to the modeler as the natural numbers have an order. However, it is the user imposing and using the ordering, not ASCEND. ASCEND sees these integers as elements in the set with NO ordering. Therefore, there are no operators in ASCEND such as successor or precursor member of a set.

Arrays

An array is a list of instances indexed over a set. The instances are all of the same *base* type (as that is the only way they can be defined). An individual member of a list may later be more refined than the other members (we shall illustrate that possibility). The following are arrays in ASCEND.

```
stage[1..n_stages]
y[components]
column[areas][processes]
```

where `components`, `areas` and `processes` are sets. For example `components` could be the set of symbols `['ethylene', 'propylene']`, `areas` the set of symbols `['feed_prep', 'prod_purification']` while `processes` could be the set `['alcohol_manuf',`

'poly\_propropylene\_manuf' ]. Note that the third example (column) is a list of lists (the way that ASCEND permits a multiply subscripted array).

The following are elements in the above arrays:

```
stage[1]
y['ethylene']
column['feed_prep']['alcohol_manuf']
```

provided that n\_stages is 1 or larger.

There can be any number of subscripts for an array. We point out, however, that in virtually every application of arrays requiring more than two subscripts, there is usually a some underlying concept that is much better modeled as an object than as part of a deeply subscripted array. In the following jagged array example, there are really the concepts of unit operation and stream that would be better understood if made explicit.

### Arrays can be jagged

(\* 4 \*) Arrays can be 'sparse' or jagged. For example:

```
process[1..3] IS_A set OF integer;
process[1] ::= [2];
process[2] ::= [7,5,3];
process[3] ::= [4,6];
FOR i in [1..3] CREATE
  FOR j IN process[i] CREATE
    flow[i][j] IS_A mass;
  END FOR;
END FOR;
```

flow is an array with six elements spread over three rows. Sparse arrays of models and variables are new to ASCEND IV.

### Arrays are also instances

Each array is itself an object. That is, when you write "a[1..2] IS\_A real;" three objects get created: a[1], a[2], and a. a is an array instance which has parts named [1] and [2] that are real instances. When a parameterized model requires an array, you pass it the single item a, not the elements a[1..2].

### Index variable

One can introduce a variable as an index ranging over a set. Index variables are local to the statements in which they occur. An example of using an index variable is the following FOR statement:

```
FOR i IN components CREATE
  VLE_equil[i]: y[i] = K[i]*x[i];
END FOR;
```

In this example *i* implicitly is of the same type as the values in the set *components*. If another object *i* exists in the model containing the FOR loop, it is ignored while executing the statements in that loop. This may cause unexpected results and the compiler will generate warnings about loop index shadowed variables.

### Label:

One can label statements which define arithmetic relationships (objective functions, equalities, and inequalities) in ASCEND. Labeling is highly recommended because it makes models much more readable and more easily debugged. Labels are also necessary for relations which are going to be used in conditional modeling or differentiation functions. A label is a sequence of alphanumeric characters ending in a colon. An example of a labeled equation is:

```
mass_balance: m_in = m_out;
```

An example of a labeled objective function is:

```
obj1: MAXIMIZE revenue - cost;
```

If a relation is defined within a FOR statement, it must have an array indexed label so that each instance created using the statement is distinguishable from the others. An example is:

```
FOR i IN components CREATE
  equil[i]: y[i] = K[i]*x[i];
END FOR;
```

The ASCEND interactive user interface identifies relationships by their labels. If one has not provided such a label, the system generates the label:

```
modelname_equationnumber
```

where *modelname* and *equationnumber* are the name of the model and the equation number in the model. An example is

```
mixture_14
```

for the unlabeled 14<sup>th</sup> relation in the mixture definition. If there is a conflict caused with an existing name, the generated name has



enough letters added after *equationnumber* to make it a unique name. Remember that each model in a refinement hierarchy inherits the equations of its less refined ancestors, so the first equation appearing in the source code of a refining model may actually be the  $n^{\text{th}}$  relation in that model.

### Lists

Often in a statement one can include a list of names or expression. A name list is one or more names where multiple list entries are separated from each other by commas. Examples of a list of names are:

```
T1, inlet_T, outlet_T
y[components], y_in
stage[1..n_stages]
```

### Ordered lists:

If the ordering of names in a list matters, that list is enclosed in (). Order matters in: calling externally defined methods or models, calling most real-valued functions, passing parameters to ASCEND models or methods, and declaring the controlling parameters that SELECT, SWITCH, and WHEN statements make decisions on.

## 20.1.3 BASIC CONCEPTS

### Instances and types

This is an opportune time to emphasize the distinction between the terms *instance* and *type*. A *type* in ASCEND is what we define when we declare an ASCEND model or atom. It is the formal definition of the attributes (parts) and attribute default values that an object will have if it is created using the type definition. Methods are associated with types.

In ASCEND there are two meanings (closely related) of an instance.

- An *instance* is a *named part* that exists within a type definition.
- An *instance* is a compiled object.

If one is in the context of the ASCEND interface, the system compiles an instance of a model type to create an object with which one carries out computations. The system requires the user to give a simple name for this simulation instance. This name given is then the first part of the qualified name for all the parts of the compiled object.

Implicit types

It is possible to create an instance that does not have a corresponding type definition in the library. The type of such an instance is said to be *implicit*. (Some people use the word *anonymous*. However, no computable type is anonymous and the implicit type of an instance is theoretically computable). The simplest example of an implicit type is the type of an instance compiled from the built-in definition `integer_constant`. For example:

```
i, j IS_A integer_constant;
i ::= 2;
j ::= 3;
```

Instances `i` and `j`, though of the same formal type, are implicit type incompatible because they have been assigned distinct values.

Instances which are either formally or implicitly type incompatible cannot be merged. This will be discussed further in Section 20.3.

Parsing

Most errors in the declaration of an ASCEND model can be caught at parse time because the object type of any well-formed name in an ASCEND definition can be resolved or proved ambiguous. We cannot prove at parse time whether a specific array element will exist, but we can know that should such an element exist, it must be of the type with which the array is defined.

Ambiguity is warned about loudly because it is caused by either misspelling or poor modeling style. The simplest example of ambiguity follows.

Assume a type, `stream`, and a refinement of `stream`, `heat_stream`, which adds the new variable `H`. Now, if we write:

```
MODEL mixer;
input[1..2] IS_A stream;
output IS_A heat_stream;
input[1].H + input[2].H = output.H;
END mixer;
```

We see the parser can find the definition of `H` in the type `heat_stream`, so `output.H` is well defined. The author of the `mixer` model may intend to refine `input[1]` and `input[2]` to be objects of different types, say `steam_stream` and `electric_stream`, where each defines an `H` suitable for use in the equation. The parser cannot read the author's mind, so it warns that `input[1].H` and `input[2].H` are ambiguous in the `mixer`

definition. The mixer model is not highly reusable except by the author, but sometimes reusability is not a high priority objective. The mixer definition is allowed, but it may cause problems in instantiation if the author has forgotten the assumption that is not explicitly stated in the model and neglects to refine the input streams appropriately.

### Instantiation

Creating an simulation based on a type definition is a multi-phase process called compiling (or instantiation). When an instantiation cannot be completed because some structural parameter (a `symbol_constant`, `real_constant`, `integer_constant`, or `set`) does not have a value there will be PENDING statements. The user interface will warn that something is incomplete.

In phase 1 all statements that create instance structures or assign constant values are executed. This phase theoretically requires an infinite number of passes through the structural statements of a definition. We allow a maximum of 5 and have never needed more than 3. There may be pending statements at the end of phase 1. The compiler or interface will issue warnings about pending statements, starting with warnings about unassigned constants.

Phase 2 compiles as many real arithmetic relation definitions as possible. Some relations may be impossible to compile because the constants or sets they depend on do not have values assigned. Other relations may be impossible because they reference variables that do not exist. This is determined in a single pass.

Phase 3 compiles as many logical arithmetic relation definitions as possible. Some relations may be impossible to compile because the constants or sets they depend on do not have values assigned. Other relations may be impossible because they reference real arithmetic relations that do not exist. This is determined in a single pass.

Phase 4 compiles as many conditional programming statements (WHENs) as possible. Some WHEN relations may be impossible to compile because the discrete variables, models, or relations they depend on do not exist. This is determined in a single pass.

Phase 5 executes the variable defaulting statements made in the declarative section of each model IF AND ONLY IF there are no pending statements from phases 1-4 anywhere in the simulation.

The first occurrence of each impossible statement will be explained during a failed compilation. Impossible statements include:

- Relations containing undefinable variables (often misspellings).
- Assignments that are dimensionally inconsistent or containing mismatched types.
- Structure building or modifying statements that refer to model parts which cannot exist or that require a type-incompatible argument, refinement, or merge.

## 20.2 DATA TYPE DECLARATIONS

In the spectrum of OO languages, ASCEND is best considered as being class-based, though it is rather more a hybrid. We have atom and model definitions, called *types*, and the compiled objects themselves, called *instances*. ASCEND instances have a record of what type they were constructed from.

### Type qualifiers:

#### **UNIVERSAL**

Universal is an optional modifier of all ATOM, CONSTANT, and MODEL definitions. If UNIVERSAL precedes the definition, then ALL instances of that type will actually refer to the first instance of the type that is created. This saves memory and ensures global consistency of data.

Examples of universal type definitions are

```
UNIVERSAL MODEL methane REFINES
generic_component_model;
```

```
UNIVERSAL CONSTANT circle_constant REFINES
real_constant ::= 1{PI};
```

```
UNIVERSAL ATOM counter_1 REFINES integer;
```

Tip: Don't use UNIVERSAL variables in relations.

It is important to note that, because variables must store information about which relations they occur in, it is a very bad idea to use UNIVERSAL typed variables in relations. The construction and maintenance of the relation list becomes very expensive for universal variables. UNIVERSAL constants are alright to use, though, because there are no relation links for constants.

## 20.2.1 MODELS

### MODEL

An ASCEND model has a declarative part and an optional procedural part headed by the METHODS word. Models are essentially containers for variables and relations. We will explain the various statements that can be made within models in Section 20.3 and Section 20.4.

### Simple models:

#### foo

```
MODEL foo;
    (* statements about foo go here*)
METHODS
    (* METHODS for foo go here*)
END foo;
```

#### bar

```
MODEL bar REFINES foo;
    (*additional statements about foo *)
METHODS
    (* additional METHODS for bar *)
END bar;
```

### Parameterized Models

(\* 4 \*) Parameterizing models makes them easier to understand and faster for the system to compile. The syntax for a parameterized model vaguely resembles a function call in imperative languages, but it is NOT. When constructing a reusable model, all the constants that determine the sizes of arrays and other structures should be declared in the parameter list so that

- the user knows what is required to reuse the model.
- the compiler knows what values must be set before it should bother attempting to compile the model.

There is no reason that other items could not also go in the parameter list, such as key variables which might be considered inputs or outputs or control parameters in the mathematical application of the model. A simple example of parameterization would be:

#### column(n,s)

```
MODEL column(
ntrays WILL_BE integer_constant;
components IS_A set of symbol_constant;
);
    stage[1..ntrays] IS_A simple_tray;
END column;
```

**flowsheet**

```

MODEL flowsheet;
    tower4size IS_A integer_constant;
    tower4size := 22;
    ct IS_A column(tower4size,['c5','c6']);
    (* additional flowsheet statements *)
END flowsheet;

```

In this example, the column model takes the first argument, `ntrays`, by reference. That is, `ct.ntrays` is an alias for the flowsheet instance `tower4size`. `tower4size` must be compiled and assigned a value before we will attempt to compile the column model instance `ct`. The second argument is taken by value, `['c5','c6']`, and assigned to components, a column part that was declared with `IS_A` in the parameter list. There is only one name for this set, `ct.components`. Note that in the flowsheet model there is no part that is a set of `symbol_constant`.

The use of parameters in ASCEND modeling requires some thought, and we will present that set of thoughts in Section 20.5. Beginners may wish to create new models without parameters until they are comfortable using the existing parameterized library definitions. Parameters are intended to support model reuse and efficient compilation which are not issues in the very earliest phase of developing novel models.

## 20.2.2 SETS

Arrays in ASCEND, as already discussed in Section 20.1.2, are defined over sets. A set is simply an instance with a set value. The elements of sets are NOT instances or sets.

### Set Declaration:

A set is made of either `symbol_constants` or `integer_constants`, so a set object is declared in one of two ways:

```

my_integer_set IS_A set OF integer_constant;
or
my_symbol_set IS_A set OF symbol_constant;

```

### **:==**

A set is assigned a value like so:

```

my_integer_set := [1,4];

```

The RHS of such an assignment must be either the name of another set instance or an expression enclosed in square brackets and made up of only set operators, other sets, and the names of

integer\_constants or symbol\_constants. Sets can only be assigned once.

### Set Operations

#### **UNION[setlist]**

A function taken over a list of sets. The result is the set that includes all the members of all the sets in the list. Note that the result of the UNION operation is an unordered set and the argument order to the union function does not matter. The syntax is:

**+** UNION[list\_of\_sets]

A+B is shorthand for UNION[A,B]

Consider the following sets for the examples to follow.

A := [1, 2, 3, 5, 9];

B := [2, 4, 6, 8];

Then UNION[A, B] is equal to the set [1, 2, 3, 4, 5, 6, 8, 9] which equals [1..6, 8, 9] which equals [[1..9] - [7]].

#### **INTERSECTION[ ]**

INTERSECTION[list of set expressions]. Finds the intersection (and) of the sets listed.

**\*** Equivalent to INTERSECTION[list\_of\_sets].

A\*B is shorthand for INTERSECTION[A,B]

For the sets A and B defined just above, INTERSECTION[A, B] is the set [2]. The \* shorthand for intersection is NOT recommended for use except in libraries no one will look at.

#### Set difference:

One can subtract one set from another. The result is the first set less any members in the set union of the first and second set. The syntax is

**-** first\_set - second\_set

For the sets A and B defined above, the set difference A - B is the set [1, 3, 5, 9] while the set difference B - A is the set [4, 6, 8].

#### **CARD[set]**

Cardinality. Returns an integer constant value that is the number of items in the set.

#### **CHOICE[set]**

Choose one. The result of running the CHOICE function over a set is an arbitrary (but consistent: for any set instance you always get the same result) single element of that set.

Running `CHOICE[A]` gives any member from the set `A`. The result is a member, not a set. To make the result into a set, it must be enclosed in square brackets. Thus `[CHOICE[A]]` is a set with a single element arbitrarily chosen from the set `A`. Good modelers do not leave modeling decisions to the compiler; they do not use `CHOICE[]`.

To reduce a set by one element, one can use the following

```
A_less_one IS_A set OF integer;
A_less_one ::= A - [CHOICE[A]];
```

**IN**

lhs `IN` rhs can only be well explained by examples. If lhs is a simple and not previously defined name, it is created as a temporary loop index which will take on the values of the rhs set definition. If lhs is something that already exists, the result of lhs `IN` rhs is context dependent; stare at the model `set_example` below which demonstrates both `IN` and `SUCH_THAT`. If you still are not satisfied, you might examine `[[westerbergksets]]`.

**SUCH\_THAT (\* 4 \*)**

Set expressions can be rather clever. We will give a detailed example because unordered sets are unfamiliar to most people and set arithmetic is quite powerful. In this example we see arrays of sets and sparse arrays.

```
MODEL set_example;
  (* we define a sparse matrix of reaction coefficient information
  * and the species balance equations. *)
  rxns IS_A set OF integer_constant;
  rxns ::= [1..3];
  species IS_A set OF symbol_constant;
  species ::= ['A','B','C','D'];

  reactants[rxns] IS_A set OF symbol_constant; (* species in each rxn_j *)
  reactants[1] ::= ['A','B','C'];
  reactants[2] ::= ['A','C'];
  reactants[3] ::= ['A','B','D'];

  reactions[species] IS_A set OF integer_constant;
  FOR i IN species CREATE (* rxns for each species i *)
    reactions[i] ::= [j IN rxns SUCH_THAT i IN reactants[j]];
  END FOR;
  (* Define sparse stoichiometric matrix. Values of eta_ij set later.*)
  FOR j IN rxns CREATE
    FOR i IN reactants[j] CREATE
      (* eta_ij --> mole i/mole rxn j*)
```



```

        eta[i][j] IS_A real_constant;
    END FOR;
END FOR;
production[species] IS_A molar_rate;
rate[rxns] IS_A molar_rate; (* mole rxn j/time *)
FOR i IN species CREATE
gen_eqn[i]: production[i] =
    SUM[eta[i][j]*rate[j] | j IN reactions[i]];
END FOR;
END set_example;

```

"|" is shorthand for  
SUCH\_THAT.

The array eta has only 8 elements, and we defined those elements in a set for each reaction. The equation needs to know about the set of reactions for a species i, and that set is calculated automatically in the model's first FOR/CREATE statement.

|

The | symbol is the ASCEND III notation for SUCH\_THAT. We noted that "|" is often read as "for all", which is different in that "for all" makes one think of a FOR loop where the loop index is on the left of an IN operator. For example, the j loop in the SUM of gen\_eqn[i] above.

### 20.2.3 CONSTANTS

ASCEND supports real, integer, boolean and character string constants. Constants in ASCEND do not have any attributes other than their value. Constants are scalar quantities that can be assigned exactly once. Constants may only be assigned using the ::= operator and the RHS expression they are assigned from must itself be constant. Constants do not have subparts. Integer and symbol constants may be used in determining the definitions of sets.

Explicit refinements of the built-in constant types may be defined as exemplified in the description of real\_constant. Implicit type refinements may be done by instantiating an incompletely defined constant and assigning its final value.

Sets could be considered constant because they are assigned only once, however sets are described separately because they are not quite scalar quantities.

#### **real\_constant**

Real number with dimensionality. Note that the dimensionality of a real constant can be specified via the type definition without immediately defining the value, as in the following pair of definitions.

CONSTANT declaration example:

```

CONSTANT molar_weight REFINES real_constant DIMENSION
M/Q;
CONSTANT hydrogen_weight REFINES molar_weight ::=
1.004{g/mole};

```

**integer\_constant**

Integer number. Principally used in determining model structure. If appearing in equations, integers are evaluated as dimensionless reals. Typical use is inside a MODEL definition and looks like:

```

n_trays IS_A integer_constant;
n_trays ::= 50;
tray[1..n_trays] IS_A vl_equilibrium_tray;

```

**symbol\_constant**

Object with a symbol value. May be used in determining model structure.

**boolean\_constant**

Logical value. May be used in determining model structure.

Setting constants**::=**

Constant and set assignment operator.

It is suggested, but not required, that names of all types that refine the built-in constant types have names that end in `_constant`.

```
LHS_list ::= RHS;
```

Here it is required that the one or more items in the LHS be of the same constant type and that RHS is a single-valued expression made up of values, operators, and other constants. The ::= is used to make clear to both the user and the system what scalar objects are constants.

## 20.2.4 VARIABLES

There are four built-in types which may be used to construct variables: symbol, boolean, integer, and real. At this time symbol types have special restrictions. Refinements of these variable base types are defined with the ATOM statement. Atom types may declare attribute fields with types real, integer, boolean, symbol, and set. These attributes are NOT independent objects and therefore cannot be refined, merged, or put in a refinement clique (ARE\_ALIKEd).

**ATOM**

The syntax for declaring a new atom type is

```
ATOM atom_type_name REFINES variable_type
```

```

    «DIMENSION dimension_expression»
    «DEFAULT value»; (* note the ; *)
    «initial attribute assignment;»
END atom_type_name;

```

### **DEFAULT, DIMENSION, and DIMENSIONLESS**

The DIMENSION attribute is for variables whose base type is real. It is an optional field. If not defined for any atom with base type real, the dimensions will be left as undefined. Any variable which is later declared to be one of these types will be given *wild card* dimensionality (represented in the interactive display by an asterisk (\*)). The system will deduce the dimensionality from its use in the relationships in which it appears or in the declaring of default values for it, if possible.

`solver_var` is a special case of ATOM and we will say much more about it in Section 20.6.1.

```

ATOM solver_var REFINES real DEFAULT 0.5 {?};
  lower_bound IS_A real;
  upper_bound IS_A real;
  nominal     IS_A real;
  fixed       IS_A boolean;
  fixed := FALSE;
  lower_bound := -1e20 {?};
  upper_bound := 1e20 {?};
  nominal := 0.5 {?};
END solver_var;

```

The default field is also optional. If the atom has a declared dimensionality, then this value must be expressed with units which are compatible with this dimensionality. In the `solver_var` example, we see a DEFAULT value of 0.5 with the unspecified unit {?} which leaves the dimensionality wild.

### **real**

Real valued variable quantity. At present, all variables that you want to be attended to by solver tools must be refinements of the type `solver_var`. This is so that modifiable parametric values can be included in equations without treating them as variables. Strictly speaking, this is a characteristic of the solver interface and not the ASCEND language. Each tool in the total ASCEND system may have its own semantics that go beyond the ASCEND object definition language.

### **integer**

Integer valued variable quantity. We find these mighty convenient for use in certain procedural computations and as attributes of `solver_var` atoms.

### **boolean**

Truth valued variable quantity. These are principally used as flags on `solver_vars` and relations. They can also be used procedurally

and as variables in logical programming models, subject to the logical solver tool's semantics. (Compare `solver_boolean` and `boolean_var` in Section 20.6.)

### **symbol**

\*4\* Symbol valued variable quantity. We do not yet have operators for building symbols out of other symbols.

### Setting variables

**:=**

Procedural equals differs from the ordinary equals (=) in that it means the left-hand-side (LHS) variables are to be assigned the value of the right-hand-side (RHS) expression when this statement is processed. Processing happens in the last phase of compiling (INSTANTIATION on page 171) or when executing a method interactively through the ASCEND user interface. The order the system encounters these statements matters, therefore, with a later result overwriting an earlier one if both statements have the same the same LHS variable.

Note that variable assignments (also known as “defaulting statements”) written in the declarative section are executed only after an instance has been fully created. This is a frequent source of confusion and errors, therefore we recommend that you **DO NOT ASSIGN VARIABLES IN THE DECLARATIVE SECTION**.

Note that := IS NOT =.

We use an ordinary equals (=) when defining a real valued equation to state that the LHS expression is to equal the RHS expression at the solution for the model. We use == for logical equations.

### Tabular assignments

(\* 4+ \*) Assigning values en masse to arrays of variables that are defined associatively on sets without order presents a minor challenge. The solution proposed in ASCEND IV (but not yet implemented as we've not had time or significant user demand) is to allow a tabular data statement to be used to assign the elements of arrays of variables or constants. The DATA statement may be used to assign variables in the declarative or methods section of a model (though we discourage its use declaratively for variable initialization) or to assign constant arrays of any type, including sets, in the declarative section. Here are some examples:

**DATA (\* 4+ \*)**

```
MODEL tabular_ex;
lset,rset,cset IS_A set OF integer_constant;
rset ::= [1..3];
cset ::= rset - [2];
lset ::= [5,7];
a[rset][cset] IS_A real;
```

```

b[lset][cset][rset] IS_A real_constant;

(* rectangle table *)
DATA FOR a:
COLUMNS 1,3; (*order last subscript cset*)
UNITS {kg/s}, {s}; (* columnar units *)
(* give leading subscripts *)
[1] 2.8, 0.3;
[2] 2.7, 1.3;
[3] 3.3, 0.6;
END DATA;

(* 2 layer rectangle table *)
CONSTANT DATA FOR b:
COLUMNS 1..3; (* order last subscript rset *)
(* UNITS omitted, so either the user gives value in the
table or values given are DIMENSIONLESS. *)
(* ordering over [lset][cset] required *)
[5][1] 3 {m}, 2{m}, 1{m};
[5][3] 0.1, 0.2, 0.3;
[7][1] -3 {m/s}, -2{m/s}, -1{m/s};
[7][3] 4.1 {1/s}, 4.2 {1/s}, 4.3 {1/s};
END DATA;

END tabular_ex;

```

For sparse arrays of variables or constants, the COLUMNS and (possibly) UNITS keywords are omitted and the array subscripts are simply enumerated along with the values to be assigned.

## 20.2.5 RELATIONS

### Mathematical expression:

The syntax for a mathematical expression is any legal combination of variable names and arithmetic operators in the normal notation. An expression may contain any number of matched rounded parentheses, (), to clarify meaning. The following is a legal arithmetic expression:

$$y^2+(\sin(x)-\tan(z))*q$$

Each additive term in a mathematical expression (terms are separated by + or - operators) must have the same dimensionality.

An expression may contain an index variable as a part of the calculation if that index variable is over a set whose elements are of type integer. (See the FOR/CREATE and FOR/DO statements below.) An example is:

```
term[i] = a[i]*x^(i-1);
```

### Numerical relations

The syntax for a numeric relation is either

```
optional_label: LHS relational_operator RHS;
or
optional_label: objective_type LHS;
```

*Objective\_type* is either MAXIMIZE or MINIMIZE. RHS and LHS must be one or more variables, constants, and operators in a normal algebraic expression. The operators allowed are defined below and in Section 20.6.3. Variable integers, booleans, and symbols are not allowed as operands in numerical relations, nor are boolean constants. Integer indices declared in FOR/CREATE loops are allowed in relations, and they are treated as integer constants.

### Relational operators:

=, >=, <=, <, >, <>

These are the numerical relational operators for declarative use.

```
Ftot*y['methane'] = m['methane'];
y['ethanol'] >= 0;
```

Equations must be dimensionally correct.

**MAXIMIZE,**  
**MINIMIZE**

Objective function indicators.

### Binary Operators:

+, -, \*, /, ^. We follow the usual algebraic order of operations for binary operators.

+

Plus. Numerical addition or set union.

-

Minus. Numerical subtraction or set difference.

\*

Times. Numerical multiplication or set intersection.

/

Divide. Numeric division. In most cases it implies real division and not integer division.

**^** Power. Numeric exponentiation. If the value of  $y$  in  $x^y$  is not integer, then  $x$  must be greater than 0.0 and dimensionless.

Unary Operators: `-, ordered_function()`

**-** Unary minus. Numeric negation. There is no unary + operator.

***ordered\_function()*** unary real valued functions. The unary real functions we support are given in section Section 20.6.3.

Real functions of sets of real terms:

**SUM[term set]** Add all expressions in the function's list.

For the SUM, the base type real items can be arbitrary arithmetic expressions. The resulting items must all be dimensionally compatible.

An examples of the use is:

```
SUM[y[components]] = 1;
```

or, equivalently, one could write:

```
SUM[y[i] | i IN components] = 1;
```

Empty SUM[] yields wild 0.

When a SUM is compiled over a list which is empty it generates a wild dimensioned 0. This will sometimes cause our dimension checking routines to fail. The best way to prevent this is to make sure the SUM never actually encounters an empty list. For example:

```
SUM[Q[possibly_empty_set], 0{watt}];
```

In the above, the variables  $Q[i]$  (if they exist) have the dimensionality associated with an energy rate. When the set is empty, the 0 is the only term in the SUM and establishes the dimensionality of the result. When the set is NOT empty the compiler will simplify away the *trailing* 0 in the sum.

**PROD[term set]** Multiply all the expressions in the product's list. The product of an empty list is a dimensionless value, 1.0.

Possible future functions: **(Not implemented - only under confused consideration at this time.)** The following functions only work in methods as they are

not smooth function and would destroy a Newton-based solution algorithm if used in defining a model equation:

**MAX[term set]** (\* 4+ \*) maximum value on list of arguments

**MIN[term set]** (\* 4+ \*) minimum value on list of arguments

### 20.2.6 DERIVATIVES IN RELATIONS (\* 4+ \*)

Simply put, we would like to have general partial and full derivatives usable in writing equations, as there are many mathematically interesting things that can be said about both. We have not implemented such things yet for lack of time and because with several implementations (see gPROMS and OMOLA, among others) already out there we can't see too many research points to be gained by more derivative work.

### 20.2.7 EXTERNAL RELATIONS

We cannot document these at the present time. The only reference for them is [[abbottthesis]].

### 20.2.8 CONDITIONAL RELATIONS (\* 4 \*)

The syntax is `CONDITIONAL list_of_relation_statements END CONDITIONAL;`

A `CONDITIONAL` statement can appear anywhere in the declarative portion of the model and it contains only relations to be used as boundaries. That is, these real arithmetic equations are used in expressing logical condition equations via the `SATISFIED` operator. See `LOGICAL FUNCTIONS` on page 206.

### 20.2.9 LOGICAL RELATIONS (\* 4 \*)

#### Logical expression

An expression whose value is `TRUE` or `FALSE` is a logical expression. Such expressions may contain boolean variables. If `A`, `B`, and `laminar` are `boolean`, then the following is a logical expression:

`A + (B * laminar)`

as is (and probably more clearly)



A OR (B AND laminar)

The plus operator acts like an OR among the terms while the times operator acts like an AND. Think of TRUE being equal to 1 and FALSE being equal to 0 with the  $1+1=0+1=1+0=1$ ,  $0+0=0$ ,  $1*1=1$  and  $0*1=1*0=0*0=0$ . If  $A = FALSE$ ,  $B=TRUE$  and  $laminar$  is TRUE, this expression has the value

FALSE OR (TRUE AND TRUE) -->TRUE

or in terms of ones and zeros

$0 + (1 * 1) --> 1.$

Logical relations are then made by putting together logical expressions with the boolean relational operators == and !=. Since we have no logical solving engine we have not pushed the implementation of logical relations very hard yet.

### 20.2.10 NOTES (\* 4+ \*)

NOTES are discussed in Chapter 5 of Allan's thesis. More details are not available here at this time as the implementation is only partially complete.

## 20.3 DECLARATIVE STATEMENTS

We have already seen several examples that included declarative statements. Here we will be more systematic in defining things. The statements we describe are legal within the declarative portion of an ATOM or MODEL definition. The declarative portion stops at the keyword METHODS if it is present in the definition or at the end of the definition.

### Statements

Statements in ASCEND terminate with a semicolon (;). Statements may extend over any number of lines. They may have blank lines in the middle of them. There may be several statements on a single line.

### Compound statements

Some statements in ASCEND can contain other statements as a part of them. The declarative compound statements are the ALIASES/ISA, CONDITIONAL, FOR/CREATE, SELECT/CASE, and WHEN/CASE statements. The procedural compound statements allowed only in methods are the FOR/DO, SWITCH (\* 4+ \*) and

the IF statements. Compound statements end with "END *word*;", where *word* matches the beginning of the syntax block, e.g. END FOR . and they can be nested, with some exceptions which are noted later.

CASE statements are here, finally!

(\*4\*) WHEN/CASE, CONDITIONAL, and SELECT/CASE handle modeling alternatives within a single definition. The easy way to remember the difference is that the first picks which equations to solve WHEN discrete *variables* have certain values, while the second SELECTs which statements to compile based on discrete *constants*. (\* 4+ \*) SWITCH statements handle flow of control in procedures, in a slightly more generalized form than the C language switch statement.

Type declarations are not compound statements.

MODEL and ATOM type definitions and METHOD definitions are not really compound statements because they require a name following their END word that matches the name given at the beginning of the definition. These definitions cannot be nested.

ASCEND operator synopses:

We'll start with an extremely brief synopsis of what each does and then give detailed descriptions. It is helpful to remember that an instance may have many names, even in the same scope, but each name may only be defined once.

**IS\_A**

Constructor. Calls for one or more named instances to be compiled using the type specified. (\* 4 \*) If the type is one that requires parameters, the parameters must be supplied in () following the type name.

**IS\_REFINED\_TO**

Reconstructor. Causes the already compiled instance(s) named to have their type changed to a more refined type. This causes an incremental recompilation of the instance(s). IS\_REFINED\_TO is not a redefinition of the named instances because refinement can only *add* compatible information. The instances retain all the structure that originally defined them. (\* 4 \*) If the type being refined to requires arguments, these must be supplied, even if the same arguments were required in the IS\_A of the original less refined declaration of the instance.

**ALIASES (\* 4 \*)**

Part alternate naming statement. Establishes another name for an instance at the same scope or in a child instance. The equivalent of an ALIASES in ASCEND III is to create another part with the desired name and merge it immediately via ARE\_THE\_SAME with the part being renamed, a rather expensive and unintuitive process.

<b>ALIASES/ISA (*4*)</b>	Creates an array of alternate names for a list of existing instances with some common base type and creates the set over which the elements of the array are indexed. Useful for making collections of related objects in ways the original author of the model didn't anticipate. Also useful for assembling array arguments to parameterized type definitions.
<b>WILL_BE (* 4 *)</b>	Forward declaration statement. Promises that a part with the given type will be constructed by an as yet unknown IS_A statement above the current scope. At present WILL_BE is legal only in defining parameters. Were it legal in the body of a model, compiling models would be very expensive.
<b>ARE_THE_SAME</b>	Merge. Calls for two or more instances already compiled to be merged recursively. This essentially means combining all the values in the instances into the most refined of the instances and then destroying all the extra, possibly less refined, instances. The remaining instance has its original name and also all the names of the instances destroyed during the merge.
<b>WILL_BE_THE_SAME (* 4 *)</b>	Structural condition statement restricting objects in a forward declaration. The objects passed to a parameterized type definition can be constrained to have arbitrary parts in common before the parameterized object is constructed.
<b>WILL_NOT_BE_THE_SAME (* 4 *)</b>	Structural condition statement restricting objects in a forward declaration. We apologize for the length of this key word, but we bet it is easy to remember. The objects passed to a parameterized type definition can be constrained to have arbitrary parts be distinct instances before the parameterized object is constructed. At present the constraint is only enforced when the objects are being passed.
<b>ARE_NOT_THE_SAME (* 4+ *)</b>	Cannot be merged. We believe it is useful to say that two objects cannot be merged and still represent a valid model. This is not yet implemented, however, mainly for lack of time. The implementation is simple.
<b>ARE_ALIKE</b>	Refinement clique constructor. Causes a group of instances to always be of the same formal type. Refining one of them causes a refinement of all the others. Does not propagate <i>implicit</i> type information, such as assignments to constants or part refinements made from a scope other than the scope of the formal definition.
<b>FOR/CREATE</b>	Indexed execution of other declarative statements. Required for creating arrays of relations and sparse arrays of other types.

- SELECT / CASE ( \* 4 \* )** Select a subset of statements to compile. Given the values of the specified *constants*, SELECT compiles all cases that match those values. A name cannot be defined two different ways inside the SELECT statement, but it may be defined outside the case statement and then *refined* in different ways in separate cases.
- CONDITIONAL ( \* 4 \* )** Describe bounding relations. The relations written inside a CONDITIONAL statement must all be labelled. These relations can be used to define regions in which alternate sets of equations apply using the WHEN statement.
- WHEN / CASE ( \* 4 \* )** When logical *variables* have certain values, use certain relations or model parts in defining a mathematical problem. The relations are not defined inside the WHEN statement because all the relations must be compiled regardless of which values the logical variables have at any given moment.

Reminder:

In the following detailed statement descriptions, we show keywords in capital letters. These words must appear in capital letters as shown in ASCEND statements. We show optional parts to a statement enclosed in double angle brackets (« ») and user supplied names in lower-case *italic* letters. (Remember that ASCEND treats the underscore (\_) as a letter). The user may substitute any name desired for these names. We use names that describe the kind of name the user should use.

Operators in detail:

**IS\_A**

This statement has the syntax

```
list_of_instance_names IS_A
model_name «(arguments_if_needed)»;
```

The IS\_A statement allows us to declare *instances* of a given *type* to exist within a model definition. If *type* has not been defined (loaded in the ASCEND environment) then this statement is an error and the MODEL it appears in is irreparably damaged (at least until you delete the type definitions and reload a corrected file). Similarly, if the arguments needed are not supplied or if provably incorrect arguments are supplied, the statement is in error. The construction of the instances does not occur until all the arguments satisfy the definition of *type*.

If a name is used twice in WILL\_BE/IS\_A/ALIASES statements of the same model, ASCEND will complain bitterly when the

definition is parsed. Duplicate naming is a serious error. Labels on relations share the same name space as other objects.

### **IS\_REFINED\_TO**

This statement has the syntax

```
list_of_instances IS_REFINED_TO
type_name «(arguments_if_needed)»;
```

We use this statement to change the type of each of the instances listed to the type *type\_name*. The modeler has to have defined each member on the list of instances. The *type\_name* has to be a type which refines the types of all the instances on the list.

An example of its use is as follows. First we define the parts called fl1, fl2 and fl3 which are of type flash.

```
fl1, fl2, fl3 IS_A flash;
```

Assume that there exists in the previously defined model definitions the type *adiabatic\_flash* that is a refinement of *flash*. Then we can make fl1 and fl3 into more refined types by stating:

```
fl1, fl3 IS_REFINED_TO adiabatic_flash;
```

This reconstruction does not occur until the arguments to the type satisfy the definition *type\_name*.

### **ALIASES (\* 4 \*)**

This statement has the syntax

```
list_of_instances ALIASES instance_name;
```

We use this statement to point at an already existing instance of any type other than *relation*, *logical\_relation*, or *when*. For example, say we want a flash tank model to have a variable T, the temperature of the vapor-liquid equilibrium mixture in the tank.

```
MODEL tank;
  feed, liquid, vapor IS_A stream;
  state IS_A VLE_mixture;
  T ALIASES state.T;
  liquor_temperature ALIASES T;
END tank;
```

We might also want a more descriptive name than T, so ALIASES can also be used to establish a second name at the same scope, e.g. *liquor\_temperature*.

An ALIASES statement will not be executed until the RHS instance has been created with an IS\_A. The compiler schedules ALIASES instructions appropriately and issues warnings if recursion is detected. An array of aliases, e.g.

```
b[1..n], c ALIASES a;
```

is permitted (though we can't think why anyone would want such an array), and the sets over which the array is defined must be completed before the statement is executed. So, in the example of b and c, the array b will not be created until a exists and n is assigned a value. b and c will be created at the same time since they are defined in the same statement. This suggests the following rule: if you must use an array of aliases, do not declare it in the same statement with a scalar alias.

The ALIASES RHS can be an element or portion of a larger array with the following exception. The existing RHS instance cannot be a relation or array of relations (including logical relations and whens) because of the rule in the language that a relation instance is associated with exactly one model.

#### **ALIASES/ISA (\*4\*)**

The ALIASES/IS\_A statement syntax is subject to change, though some equivalent will always exist. We take a set of *symbol\_constant* or *integer\_constant* and pair it with a list of instances to create an array. For the moment, the syntax and semantics is as follows.

```
alias_array_instance[aset] ALIASES
(list_of_instances) WHERE aset IS_A set OF
settype;
```

or

```
alias_array_instance[aset] ALIASES
(list_of_instances) WHERE aset IS_A set OF settype
WITH_VALUE (value_list_matching_settype);
```

*aset* is the name of the set that will be created by the IS\_A to index the array of aliases. If *value\_list\_matching\_set\_type* is not given, the compiler will make one up out of the integers (1..number of names in *list\_of\_instances*) or symbols derived from the individual names given. If the value list is given, it must have the same number of elements as the list of instances does. The value list elements must be unique because they form a set. The list of

instances can contain duplicates. If any of these conditions are not met properly, the statement is in error.

ALIASES/IS\_A can be used inside a FOR statement. When this occurs, the definition of *aset* must be indexed and it must be the last subscript of *alias\_array\_instance*. The statement must look like:

```
array_instance[FOR_index][aset[FORindex]]
ALIASES (list_of_instances) WHERE
aset[FORindex] IS_A set OF settype WITH_VALUE
(value_list_matching_settype);
```

Here, as with the unindexed version, the WITH\_VALUE portion is optional.

If this explanation is unclear, just try it out. The compiler error messages for ALIASES/IS\_A are particularly good because we know it is a bit tricky to explain.

#### WILL\_BE (\* 4 \*)

```
instance WILL_BE type_name;
```

The most common use of this forward declaration is as a statement within the parameter list of a model definition. In parameter lists, *list\_of\_instances* must contain exactly one instance. When a model definition includes a parameter defined by WILL\_BE, that model cannot be compiled until a compiled instance at least as refined as the type specified by *type\_name* is passed to it.

(\* 4+ \*) The second potential use of WILL\_BE is to establish that an array of a common base type exists and its elements will be filled in individually by IS\_A or ARE\_THE\_SAME or ALIASES statements. WILL\_BE allows us to avoid costly reconstruction or merge operations by establishing a placeholder instance which contains just enough type information to let us check the validity of other statements that require type compatibility while delaying construction until it is called for by the filling in statements. Instances declared with WILL\_BE are never compiled if they are not ultimately resolved to another instance created with IS\_A. Unresolved WILL\_BE instances will appear in the user interface as objects of type PENDING\_INSTANCE\_*model\_name*. Because of the many implementation and explanation difficulties this usage of WILL\_BE creates, it is not allowed. The ALIASES/IS\_A construct does the same job in a much simpler way.

#### ARE\_THE\_SAME

The format for this instruction is

*list\_of\_instances* ARE\_THE\_SAME;

All items on the list must have compatible types. For the example in Fig. 1, consider a model where we define the following parts:

```
a1 IS_A A;  
b1 IS_A B;  
c1 IS_A C;  
d1 IS_A D;  
e1 IS_A E;
```

Then the following ARE\_THE\_SAME statement is legal

```
a1, b1, c1 ARE_THE_SAME;
```

while the following are not

```
b1, d1 ARE_THE_SAME;  
a1, c1, d1 ARE_THE_SAME;  
b1, e1 ARE_THE_SAME;
```

When compiling a model, ASCEND will put all of the instances mentioned as being the same into an ARE\_THE\_SAME “clique.” ASCEND lists members of this clique when one asks via the interface for the aliases of any object in a compiled model.

Merging any other item with a member of the clique makes it the same as all the other items in the clique, i.e., it adds the newly mentioned items to the existing clique.

ASCEND merges all members of a clique by first checking that all members of the clique are type compatible. It then changes the type designation of all clique members to that of the most refined member.



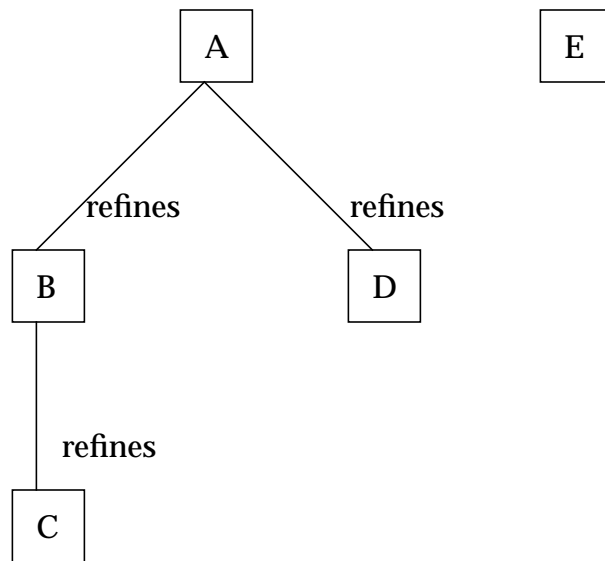


Figure 1. Diagram of the model type hierarchy A,B,C,D,E

It next looks inside each of the instances, all of which are now of the same type, and puts all of the parts with the same name into their respective ARE\_THE\_SAME cliques. The process repeats by processing these cliques until all parts of all parts of all parts, etc., are their respective most refined type or discovered to be type incompatible.

There are now lots of cliques associated with the instances being merged. The type associated with each such clique is now either a model, an array, or an atom (i.e., a variable, constant, or set). If a model, only one member of the clique generates its equations. If a variable, it assigns all members to the same storage location.

Note that the values of constants and sets are essentially *type* information, so merging two already assigned constants is only possible if merging them does not force one of them to be assigned a new value. Merging arrays with mismatching ranges of elements is an error.

**WILL\_BE\_THE\_SAME**  
 (\* 4 \*)

There is no further explanation of WILL\_BE\_THE\_SAME.

**WILL\_NOT\_BE\_THE\_SAME**  
 (\* 4 \*)

There is no further explanation of WILL\_NOT\_BE\_THE\_SAME.

**ARE\_NOT\_THE\_SAME**  
(\* 4+ \*)

ARE\_NOT\_THE\_SAME will be documented further when it is implemented.

**ARE\_ALIKE**

The format for this statement is

```
list_of_instance_names ARE_ALIKE;
```

The compiler places all instances in the list into an ARE\_ALIKE clique. It checks that the members are formally type compatible and then it converts each into the most refined type of any instance in the clique. At that point the compiler stops. It does not continue by placing the parts into cliques nor does it merge anything.

There are important consequences of modeling with such a partial merge. The consequences we are about to describe can be much more reliably achieved by use of parameterized types, *when the types are well understood*. When we are exploring new ways of modeling, ARE\_ALIKE still has its uses. When a model and its initial uses are understood well enough to be put into a reusable library, then parameterization and the explicit statement of structural constraints by operators such as WILL\_NOT\_BE\_THE\_SAME should be the preferred method of ensuring correct use.

One consequence of ARE\_ALIKE is to prevent extreme model misuse when configuring models. For example, suppose a modeler creates a new pressure changing model. The modeler is not yet concerned about the type of the streams into and out of the device but does care that these streams are of the same final type. For example, the modeler wants both to be liquid streams if either is or both to be vapor streams if either is. By declaring both to be streams only but declaring the two streams to be alike, the modeler accomplishes this intent. Suppose the modeler merges the inlet stream with a liquid outlet stream from a reactor. The merge operation makes the inlet stream into a liquid stream. The outlet stream, being in an ARE\_ALIKE clique with the inlet stream, also becomes a liquid stream. Any subsequent merge of the outlet stream with a vapor stream will lead to an error due to type incompatibility when ASCEND attempts to compile that merge. Without the ARE\_ALIKE statement, the compiler can detect no such incompatibility unless parameterized models are used.

Another purpose is the propagation of type through a model. Altering the type of the inlet stream through merging it with a liquid stream automatically made the outlet stream into a liquid stream.

If all the liquid streams within a distillation column are alike, then the modeler can make them all into streams with a particular set of components in them and with the same method used for physical property evaluation by merging only one of them with a liquid stream of this type. This is *the primary example* which has been used to justify the existence of ARE\_ALIKE. We have observed that its use makes a column library very difficult to compile efficiently. But since we now have parameterized types to help us keep the column library semantically consistent, ARE\_ALIKE can be left to its proper role: the rapid prototyping of partially understood models.

Finally, because ARE\_ALIKE does not recursively put the parts of ARE\_ALIKEd instances into ARE\_ALIKE cliques, it is possible to ARE\_ALIKE model instances which have compatible formal types but incompatible *implicit* types. This can lead to unexpected problems later and makes the ARE\_ALIKE instruction a source of non-reusability.

#### FOR/CREATE

The FOR/CREATE statement is a compound statement that looks like a loop. It isn't, however, necessarily compiled as a loop. What FOR really does is specify an index set value. Its format is:

```
FOR index_variable IN set CREATE
    list_of_statements;
END FOR;
```

This statement can be in the declarative part of the model definition only. Every statement in the list should have at least one occurrence of the index variable, or the statement should be moved outside the FOR to avoid redundant execution. A correct example is

```
FOR i IN components CREATE
    a.y[i], b[i] ARE_THE_SAME;
    y[i] = K[i]*x[i];
END FOR;
```

FOR loops can be nested to produce sparse arrays as illustrated in ARRAYS CAN BE JAGGED on page 167. IS\_A and ALIASES statements are allowed in FOR loops, provided the statements are properly indexed, a new feature in ASCEND IV.

#### SELECT/CASE (\*4\*)

Declarative. Order does not matter. All matching cases are executed. The OTHERWISE is executed if present and no other CASEs match. SELECT is not allowed inside FOR. Writing FOR statements inside SELECT is allowed.

- CONDITIONAL (\*4\*)** Both real and logical relations are allowed in **CONDITIONAL** statements. **CONDITIONAL** is really just a shorthand for setting the \$boundary flag on a whole batch of relations, since \$boundary is a write-once attribute invisible through the user interface and methods at this time.
- WHEN/CASE (\* 4 \*)** Inside each **CASE**, relations or model parts to be used are specified by writing, for example, **USE mass\_balance\_1;**. The method of dealing with the combined logical/nonlinear model is left to the solver. All matching **CASEs** are included in the problem to be solved. We do not yet have a solver which dynamically determines the applicable set of relations during solution. Our solver interface currently sends only the equations specified by the current values of the discrete variables to the client solving engine.

## 20.4 PROCEDURAL STATEMENTS

**METHODS** This statement separates the method definitions in **ASCEND** from the declarative statements. All statements following this statement are to define methods in **ASCEND** while all before it are for the declarative part of **ASCEND**. The syntax for this statement is simply

**METHODS**

with no punctuation. The next code must be a **METHOD** or the **END** of the type being defined. If there are no method definitions, this statement may be omitted.

(\* 4+ \*) **METHOD** definitions for a type can also be added or replaced after the type has been defined. This is to make creating and debugging of methods as interactive as possible. Currently, an instance must be destroyed and recreated each time a new or revised method is added to the type definition. This is a very expensive process when working with models of significant size.

The detailed semantics of method inheritance, addition, and replacement of methods are given at the end of this section.

**ADD METHODS IN type\_name; (\*4+\*)** This statement allows new methods to be added to an already loaded type definition. The next code must be a **METHOD** or the **END METHODS;** statement. If a method of the same name already exists in `type_name`, the statement is in error.

**REPLACE METHODS**  
**IN type\_name;**  
**(\*4+\*)**

This statement allows existing methods to be replaced in an already loaded type definition. The next code must be a METHOD or the END METHODS; statement. If a method of the same name does not exist in type\_name, the statement is in error.

### Initialization routines:

#### **METHOD**

A method in ASCEND must appear following the METHODS statement within a model. The system executes procedural statements of the method in the order they are written.

At present, there are no local variables or other structures in methods except loop indices. A method may be written recursively, but there is an arbitrary stack depth limit (currently set to 20 in compiler/initialize.h) to prevent the system from crashing on infinite recursions.

Specifically disallowed in ASCEND III methods are IS\_A, ALIASES, WILL\_BE, IS, IS\_REFINED\_TO, ARE\_THE\_SAME and ARE\_ALIKE statements as these “declare” the structure of the model and belong only in the declarative section.

(\* 4+ \*) In the near future, declarations of local instances (which are automatically destroyed when the method exits) will be allowed. Since methods are imperative, these local structure definitions are processed in the order they are written. Local structures are not allowed to shadow structures in the model context with which the method is called. When local structures are allowed, it will also be possible to define methods which take parameters and return values, thereby making the imperative ASCEND methods a rapid prototyping tool every bit as powerful and easy to use as the declarative ASCEND language.

The syntax for a method declaration is

```
METHOD method_name;
    «procedural statement;» (*one or more*)
END method_name;
```

### Procedural assignment

The syntax is

```
instance_name := mathematical_expression;
or
array_name[set_name] := expression;
or
list_of_instance_names := expression.
```

Its meaning is that the value for the variable(s) on the LHS is set to the value of the expression on the RHS.

DATA statements (DATA (\* 4+ \*) on page 180) can (should, rather) also appear in methods.

#### **FOR/DO statement**

This statement is similar to the FOR/CREATE statement except it can only appear in a method definition. An example would be

```
FOR i IN [1..n_stages] DO
    T[i] := T[1] + (i-1)*DT;
    ...
END FOR;
```

Here we actually execute using the values of *i* in the sequence given. So,

```
FOR i IN [n_stages..1] DO ... END FOR;
```

is an empty loop, while

```
FOR i IN [n_stages..1] DECREASING DO ... END FOR;
```

is a backward loop.

#### **IF**

The IF statement can only appear in a method definition. Its syntax is

```
IF logical_expression THEN
    list_of_statements
ELSE
    list_of_statements
END IF;
```

or

```
IF logical_expression THEN
    list_of_statements
END IF;
```

If the logical expression has a value of TRUE, ASCEND will execute the statements in the THEN part. If the value is FALSE, ASCEND executes the statements in the optional ELSE part. Please use () to make the precedence of AND, OR, NOT, ==, and != clear to both the user and the system.

- SWITCH (\* 4+ \*)** Essentially equivalent to the C switch statement, except that ASCEND allows wildcard matches, and any number of controlling variables to be given in a list.
- CALL** External calls are not presently well defined, pending debugging of the EXTERNAL connection prototype originally created by Kirk Abbott.
- RUN** This statement can appear only in a method. Its format is:

```
RUN name_of_method;
or
RUN part_name.name_of_method;
or
RUN model_type::name_of_method;
```

The named method can be defined in the current model (the first syntax), or in any of its parts (the second syntax). Methods defined in a part will be run in the scope of that part, not at the scope of the RUN statement.

- Type access to methods: When *model\_type::* appears, the type named must be a type that the current model is refined from. In this way, methods may be defined incrementally. For example:

```
MODEL foo;
  x IS_A generic_real;
METHODS
METHOD specify;
  x.fixed:= TRUE;
END specify;
END foo;

MODEL bar REFINES foo;
  y IS_A generic_real;
METHODS
METHOD specify;
  RUN foo::specify;
  y.fixed := TRUE;
END specify;
END bar;
```

## 20.5 PARAMETERIZED MODELS

Parameterized model definitions have the following general form.

```
MODEL new_type(parameter_list;)
«WHERE (where_list;)»
«REFINES existing_type «(assignment_list;)»»;
```

### 20.5.1 THE PARAMETER LIST

A parameter list is a list of statements about the objects that will be passed into the model being defined when an instance of that model is created by IS\_A or IS\_REFINED\_TO. The parameter list is designed to allow a complete statement of the necessary and sufficient conditions to construct the parameterized model. The mechanism implemented is general, however, so it is possible to put less than the necessary information in the parameter list if one seeks to confuse the model's reusers. To make parameters easy to understand for users with experience in other computer languages (and to make the implementation much simpler), we define the parameter list as ordered. All the statements in a parameter list, including the last one, must end with a ";". A parameter list looks like:

```
MODEL test (
  x WILL_BE real;
  n IS_A integer_constant;
  p[1..n] IS_A integer_constant;
  q[0..2*n-1] WILL_BE widget;
);
```

Each WILL\_BE statement corresponds to a single instance that the user must create and pass into the definition of test. We will establish the local name *x* for the first instance passed to the definition of test. *n* is handled similarly, and it must precede the definition of *p*[1..*n*], because it defines the set for the array *p*. Constant types can also be defined with WILL\_BE, though we have used IS\_A for the example test.

Each IS\_A statement corresponds to a single constant valued instance or an array of constant valued instances that we will create as part of the model we are defining. Thus, the user of test must supply an array of constants as the third argument. We will check that the instance supplied is subscripted on the set [1..*n*] and copy the corresponding values to the array *p* we create local to the instance of test.



WILL\_BE statements can be used to pass complex objects (models) or arrays of objects. Both WILL\_BE and IS\_A statements can be passed arguments that are *more* refined than the type listed. If an object that is *less* refined than the type listed, the instance of parameterized model test will not be compiled. When a parameterized model type is specified with a WILL\_BE statement, NO arguments should be given. We are only interested in the formal type of the argument, not how it was constructed.

### 20.5.2 THE WHERE LIST

We can write structural and equation constraints on the arguments in the WHERE list. Each statement is a WILL\_BE\_THE\_SAME, a WILL\_NOT\_BE\_THE\_SAME, an equation written in terms of sets or discrete constants, or a FOR/CREATE statement surrounding a group of such statements. Until all the conditions in the WHERE list are satisfied, an object cannot be constructed using the parameterized definition. If the arguments given to a parameterized type in an IS\_A or IS\_REFINED\_TO statement cannot possibly satisfy the conditions, the IS\_A or IS\_REFINED\_TO statement is abandoned by the compiler.

We have not created a WILL\_BE\_ALIKE statement because formal type compatibility in ASCEND is not really a meaningful guarantee of object compatibility. Object compatibility is much more reliably guaranteed by checking conditions on the structure determining constants of a model instance.

### 20.5.3 THE ASSIGNMENT LIST

When we declare constant parameters with IS\_A, we can in a later refinement of the parameterized model assign their values in the assignment list, thus removing them from the parameter list. If an array of constants is declared with IS\_A, then we must assign values to ALL the array elements at the same time if we are going to remove them from the parameter list. If an array element is left out, the type which assigns some of the elements and any subsequent refinements of that type will not be compilable.

### 20.5.4 REFINING PARAMETERIZED TYPES

Because we wish to make the parameterized model lists represent all the parameters and conditions necessary to use a model of any

type, we must repeat the parameters declared in the ancestral type when we make a refinement. If we did not repeat the parameters, the user would be forced to hunt up the (possibly long) chain of types that yield an interesting definition in order to know the list of parameters and conditions that must be satisfied in order to use a model. We repeat all the parameters of the type being refined before we add new ones. The only exception to this is that parameters defined with `IS_A` and then assigned in the `assignment_list` are not repeated because the user no longer needs to supply these values. A refinement of the model `test` given in Section 20.5.1 follows.

```
MODEL expanded_test (
  x WILL_BE real;
  p[1..n] IS_A integer_constant;
  q[0..2*n-1] WILL_BE better_widget;
  r[0..q[0].k] WILL_BE gizmo;
  ms WILL_BE set OF symbol_constant;
) WHERE (
  q[0].k >= 2;
  r[0..q[0].k].giz_part WILL_BE THE SAME;
) REFINES test(
  n ::= 4;
);
```

In `expanded_test`, we see that the type of the array `q` is more refined than it was in `test`. We see that constants and sets from inside passed objects, such as `q[0].k`, can be used to set the sizes of subsequent array arguments. We see a structural constraint that all the gizmos in the array `r` must have been constructed with the same `giz_part`. This condition probably indicates that the gizmo definition takes `giz_part` as a `WILL_BE` defined parameter.

## 20.6 MISCELLANY

### 20.6.1 VARIABLES FOR SOLVERS

#### **solver\_var**

`Solver_var` is the base-type for all *computable* variables in the current ASCEND system. Any instances of an atom definition that refines `solver_var` are considered potential variables when constructing a problem for one of the solvers.

`Solver_var` has wild card dimensionality. (Wild card means that until ASCEND can decide what its dimensionality is, it has none

assigned. ASCEND can decide on dimensionality while compiling or executing.) In system.lib we define the following parts with associated initial values for each:

<u>Attributes:</u>	type	default
<b>lower_bound</b>	real	0.0
<b>upper_bound</b>	real	0.0
<b>nominal</b>	real	0.0
<b>fixed</b>	boolean	FALS E

*lower\_bound* and *upper\_bound* are bounds for a variable which are monitored and maintained during solving. The nominal value the value used to scale a variable when solving. The flag *fixed* indicates if the variable is to be held fixed during solving. All atoms which are refinements of *solver\_var* will have these parts. The refining definitions may reassign the default values of the attributes.

The latest full definition of *solver\_var* is always in the file system.lib.

### **generic\_real**

**One should not declare a variable to be of type *solver\_var*.** The nominal value and bound values will get you into trouble when solving. If you are programming and do not wish to declare variable types, then declare them to be of type *generic\_real*. This type has nominal value of 0.5 and lower and upper bounds of -1.0e50 and 1.0e50 respectively. It is dimensionless. *Generic\_real* is the first refinement of *solver\_var* and is also defined in system.lib

### Kluges for MILPs

Also defined in system.lib are the types for integer, binary, and semi-continuous variables.

### **solver\_semi, solver\_integer, solver\_binary**

We define basic refinements of *solver\_var* to support solvers which are more than simply algebraic. Various mixed integer-linear program solvers can be fed *solver\_semi* based atoms defining semi-continuous variables, *solver\_integer* based atoms defining integer variables, and *solver\_binary* based atoms defining binary variables.

Integers are relaxable.

All these types have associated boolean flags which indicate that either the variable is to be treated according to its restricted meaning or it is to be relaxed and treated as a normal algebraic variable.

Kluges for ODEs

We have an alternate version of `system.lib` called `ivpsystem.lib` which adds extra flags to the definition of `solver_var` in order to support initial value problem (IVP) solvers (integrators). Integration in the ASCEND IV environment is explained in another chapter.

**ivpsystem.lib**

Having `ivpsystem.lib` is a temporary, but highly effective, way to keep people who want to use ASCEND only for algebraic purposes from having to pay for the IVP overhead. Algebraic users load `system.lib`. Users who want both algebraic and IVP capability load `ivpsystem.lib` instead of `system.lib`. This method is temporary because part of the extended definition of ASCEND IV is that differential calculus constructs will be explicitly supported by the compiler. The calculus is not yet implemented, however.

**20.6.2 SUPPORTED ATTRIBUTES**

(\* 4+ \*)

The `solver_var`, and in fact most objects in ASCEND IV, should have built-in support for (and thereby efficient storage of) quite a few more attributes than are defined above. These built-in attributes are not instances of any sort, merely values. The syntax for naming one of these supported attributes is:

*object\_name* . *\$supported\_attribute\_name*.

Supported attributes may have symbol, real, integer, or boolean values. Note that the `$` syntax is essentially the same as the derivative syntax for relations; derivatives are a supported attribute of relations. The supported attributes must be defined at the time the ASCEND compiler is built. The storage requirement for a supported boolean attribute is 1 bit rather than the 24 bytes required to store a run time defined boolean flag. Similarly, the requirement for a supported real attribute is 4 or 8 bytes instead of 24 bytes.

**20.6.3 SINGLE OPERAND REAL FUNCTIONS:**

<b>exp()</b>	exponential (i.e., $\exp(x) = e^x$ )
<b>ln()</b>	log to the base e
<b>sin()</b>	sine. argument must be an angle.
<b>cos()</b>	cosine. argument must be an angle.
<b>tan()</b>	tangent. argument must be an angle.

<b>arcsin()</b>	inverse sine. return value is an angle.
<b>arccos()</b>	inverse cosine. return value is an angle.
<b>arctan()</b>	inverse tangent. return value is an angle.
<b>erf()</b>	error function
<b>sinh()</b>	hyperbolic sine
<b>cosh()</b>	hyperbolic cosine
<b>tanh()</b>	hyperbolic tangent
<b>arcsinh()</b>	inverse hyperbolic sine
<b>arccosh()</b>	inverse hyperbolic cosine
<b>arctanh()</b>	inverse hyperbolic tangent
<b>lnm()</b>	modified ln function. This lnm function is parameterized by a constant a, which is typically set to about 1.e-8. lnm(x) is defined as follows:

$\ln(x)$  for  $x > a$

$(x-a)/a + \ln(a)$  for  $x \leq a$ .

Below the value a (default setting is 1.0e-8), lnm takes on the value given by the straight line passing through  $\ln(a)$  and having the same slope as  $\ln(a)$  has at a. This function and its first derivative are continuous. The second derivative contains a jump at a.

The lnm function can tolerate a negative argument while the ln function cannot. At present the value of a is controllable via the user interface of the ASCEND solvers.

Operand dimensionality must be correct.

The operands for an ASCEND function must be dimensionally consistent with the function in question. Most transcendental functions require dimensionless arguments. The trigonometric functions require arguments with dimensionality of plane angles, P. ASCEND functions return dimensionally correct results.

The operands for ASCEND functions are enclosed within rounded parentheses, (). An example of use is:

$$y = A * \exp(-B/T);$$

Discontinuous functions: Discontinuous functions may destroy a Newton-based solution algorithm if used in defining a model equation. We strongly suggest considering alternative formulations of your equations.

**abs ( )** absolute value of argument. Any dimensionality is allowed in an abs() function.

## 20.6.4 LOGICAL FUNCTIONS

**SATISFIED( ) (\*4\*)** SATISFIED(relation\_name,tolerance) returns TRUE if the relation named has a residual value less than the real value, tolerance, given. If the relation named is a logical relation, the tolerance should not be specified, since logical relations evaluate directly to TRUE or FALSE.

## 20.6.5 UNITS DEFINITIONS

As noted in 20.1.2, ASCEND will recognize conversion factors when it sees them as {units). These units are built up from the basic units, and new units can be defined by the user. Note that the assignment `x:= 0.5 {100}`; yields `x == 50`, and that there are no 'offset conversions,' e.g. `F=9/5C+32`. Please keep unit names to 20 characters or less as this makes life pretty for other users

One or more unit conversion factors can be defined with the UNITS keyword. A unit of measure, once defined, stays in the system until the system is shut down. A measuring unit cannot be defined differently without first shutting down the system, but duplicate or equivalent definitions are quietly ignored.

A UNITS declaration can occur in a file by itself, inside a model or inside an atom. UNITS definitions are parsed immediately, they will be processed even if a surrounding MODEL or ATOM definition is rejected. Because units and dimensionality are designed into the deepest levels of the system, a unit definition must be parsed before any atoms or relations use that definition. It is good design practice to keep customized unit definitions in separate files and REQUIRE those files at the beginning of any file that uses them. Unit definitions are made in the form, for example:

```
UNITS (* several unit definitions could be here. *)
    ohm = {kilogram*meter^2/second^3/ampere^2};
END UNITS;
```

The standard units library, measures.a4l, is documented in Chapter 21.

























# CHAPTER 21 UNITS LIBRARY

## 21.1 UNITS

This chapter defines the dimensions and units and all the attendant conversion factors. Note that all conversions are simply multiplicative. This information is from the file `models/measures.a4l` in the ASCEND source code.

Note that units can be easily defined to suit the needs of local users. We are always on the lookout for new and interesting units, so if you have some send them in. From `measures.a4l` we have:

## 21.2 THE BASIC UNITS IN AN EXTENDED SI MKS SYSTEM

These units (kilogram, mole, et c.) are associated with the dimensionality listed here (M, Q, et c.) by the ASCEND IV C code. All other units are derived from these by multiplication factors. Use of units other than these requires loading unit definitions, either from `measures.a4l` or from another ASCEND file containing a UNITS declaration. The system rejects loudly any model or variable definition using undefined units.

```
define kilogram    M; (* internal mass unit SI *)
define mole        Q; (* internal quantity unit SI *)
define second      T;  (* internal time unit SI *)
define meter       L;  (* internal length unit SI *)
define Kelvin      TMP; (* internal temperature unit SI *)
define currency    C;  (* internal currency unit *)
define ampere      E;  (* internal electric current unit SI suggested *)
```

```

define candela    LUM; (* internal luminous intensity unit SI *)
define radian     P;  (* internal plane angle unit SI suggested *)
define steradian  S;  (* internal solid angle unit SI suggested *)

```

### 21.3 UNITS DEFINED IN MEASURES.A4L, THE DEFAULT SYSTEM UNITS LIBRARY OF ATOMS.A4L.

```

distance      pc = 3.08374e+16*meter;

                parsec = pc;

                kpc = 1000*pc;

                Mpc = 1e6*pc;

                km = meter*1000;

                m = meter;

                dm = meter/10;

                cm = meter/100;

                mm = meter/1000;

                um = meter/1000000;

                nm = 1.e-9*meter;

                kilometer = km;

                centimeter = cm;

                millimeter = mm;

                micron=um;

                nanometer = nm;

                angstrom = m/1e10;

                fermi = m/1e15;

```

mi = 1609.344\*meter;  
yd = 0.914412\*meter;  
ft = 0.304804\*meter;  
inch = 0.0254\*meter;  
mile = mi;  
yard = yd;  
feet = ft;  
foot = ft;  
in = inch;  
mass metton = kilogram \*1000;  
mton = kilogram \*1000;  
kg = kilogram;  
g = kilogram/1000;  
gram = g;  
mg = g/1000;  
milligram = mg;  
ug= kilogram\*1e-9;  
microgram = ug;  
ng=kilogram\* 1e-12;  
nanogram=ng;  
pg=kilogram\* 1e-15;  
picogram=pg;  
amu = 1.661e-27\*kilogram;

time

lbm = 4.535924e-1\*kilogram;

ton = lbm\*2000;

oz = 0.028349525\*kilogram;

slug = 14.5939\*kilogram;

yr = 31557600\*second;

wk = 604800\*second;

dy = 86400\*second;

hr = 3600\*second;

min = 60\*second;

sec = second;

s = second;

ms = second/1000;

us = second/1e6;

ns = second/1e9;

ps = second/1e12;

year = yr;

week = wk;

day = dy;

hour = hr;

minute = min;

millisecond = ms;

microsecond = us;

nanosecond = ns;

picosecond = ps;

molecular quantities kg\_mole=1000\*mole;

g\_mole = mole;

gm\_mole = mole;

kmol = 1000\*mole;

mol = mole;

mmol = mole/1000;

millimole=mmol;

umol = mole/1e6;

micromole=umol;

lb\_mole = 4.535924e+2\*mole;

temperature K = Kelvin;

R = 5\*Kelvin/9;

Rankine = R;

money dollar = currency;

US = currency;

USDollar=currency;

CR = currency;

credits=currency;

reciprocal time  
(frequency) rev = 1.0;

cycle = rev;

rpm = rev/minute;

rps = rev/second;

hertz = cycle/second;

Hz = hertz;

area

ha = meter<sup>2</sup>\*10000;

hectare=ha;

acre= meter<sup>2</sup>\*4046.856;

volume

l = meter<sup>3</sup>/1000;

liter = l;

ml = liter/1000;

ul = liter/1e6;

milliliter = ml;

microliter = ul;

hogshead=2.384809e-1\*meter<sup>3</sup>;

cuft = 0.02831698\*meter<sup>3</sup>;

impgal = 4.52837e-3\*meter<sup>3</sup>;

gal = 3.785412e-3\*meter<sup>3</sup>;

barrel = 42.0\*gal;

gallon = gal;

quart = gal/4;

pint = gal/8;

cup = gal/16;

floz = gal/128;

force

N = kilogram\*meter/second<sup>2</sup>;

newton = N;

dyne = N\*1.0e-5;

pn=N\*1e-9;

picoNewton=pn;

lbf = N\*4.448221;

pressure Pa = kilogram/meter/second^2;

MPa = 1.0e+6\*Pa;

bar =1.0e+5\*Pa;

kPa = 1000\*Pa;

pascal = Pa;

atm = Pa\*101325.0;

mmHg = 133.322\*Pa;

torr = 133.322\*Pa;

psia = 6894.733\*Pa;

psi = psia;

ftH2O = 2989\*Pa;

energy J = kilogram\*meter^2/second^2;

joule = J;

MJ = J \* 1000000;

kJ = J \* 1000;

mJ=J\*1.0e-3;

uJ=J\*1.0e-6;

nJ=J\*1.0e-9;

milliJoule=mJ;



power

microJoule=uJ;  
nanoJoule=nJ;  
erg = J\*1.0e-7;  
BTU = 1055.056\*J;  
pCu = BTU \* 1.8;  
cal = J\*4.18393;  
calorie = cal;  
kcal=1000\*calorie;  
Cal=1000\*calorie;  
W = J/second;  
EW = 1.0e+18\*W;  
PW = 1.0e+15\*W;  
TW = 1.0e+12\*W;  
GW = 1.0e+9\*W;  
MW = 1.0e+6\*W;  
kW = 1000\*W;  
mW = W/1000;  
uW = W/1000000;  
nW = W/1e9;  
pW = W/1e12;  
fW = W/1e15;  
aW = W/1e18;  
terawatt = TW;

gigawatt = GW;

megawatt = MW;

kilowatt = kW;

watt = W;

milliwatt = mW;

microwatt = uW;

nanowatt = nW;

picowatt = pW;

femtowatt = fW;

attowatt = aW;

hp= 7.456998e+2\*W;

absolute viscosity

poise = Pa\*s;

cP = poise/100;

electric charge

coulomb=ampere\*second;

C = coulomb;

coul = coulomb;

mC = 0.001\*C;

uC = 1e-6\*C;

nC = 1e-9\*C;

pC = 1e-12\*C;

miscellaneous electro-  
magnetic fun

V = kilogram\*meter^2/second^3/ampere;

F = ampere^2\*second^4/kilogram/meter^2;

ohm = kilogram\*meter^2/second^3/ampere^2;

$$\text{mho} = \text{ampere}^2 * \text{second}^3 / \text{kilogram} / \text{meter}^2;$$

$$S = \text{mho};$$

$$\text{siemens} = S;$$

$$A = \text{ampere};$$

$$\text{amp} = \text{ampere};$$

$$\text{volt} = V;$$

$$\text{farad} = F;$$

$$\text{mA} = A / 1000;$$

$$\text{uA} = A / 1000000;$$

$$\text{kV} = 1000 * V;$$

$$\text{MV} = 1e6 * V;$$

$$\text{mV} = V / 1000;$$

$$\text{mF} = 0.001 * F;$$

$$\text{uF} = 1e-6 * F;$$

$$\text{nF} = 1e-9 * F;$$

$$\text{pF} = 1e-12 * F;$$

$$\text{kohm} = 1000 * \text{ohm};$$

$$\text{Mohm} = 1e6 * \text{ohm};$$

$$\text{kS} = 1000 * S;$$

$$\text{mS} = 0.001 * S;$$

$$\text{uS} = 1e-6 * S;$$

$$\text{Wb} = V * \text{second};$$

$$\text{weber} = \text{Wb};$$

tesla = Wb/m<sup>2</sup>;

gauss = 1e-4\*tesla;

H = Wb/A;

henry = H;

mH = 0.001\*H;

uH = 1e-6\*H;

numeric constants of  
some interest

To set a variable or constant to these, the code is (in the declarations):

```

ATOM unspecified_unitwise REFINES real;
END unspecified_unitwise;
MODEL gizmo;
  x IS_A unspecified_unitwise;
  (* if some other atom type is more appropriate, by all
  * means, use it.
  *)
  x := 1 {PI};
  ...
END gizmo;

molecule = 1.0;
PI=3.141592653589793;           # Circumference/Diameter ratio
EULER_C = 0.57721566490153286;  # euler gamma
GOLDEN_C = 1.618033988749894;   # golden ratio
HBAR = 1.055e-34*J*second;      # Reduced Planck's constant
PLANCK_C = 2*PI*HBAR;          # Planck's constant
LIGHT_C = 2.99793e8 * meter/second; # Speed of light in vacuum
MU0 = 4e-7*PI*kg*m/(C*C);      # Permeability of free space
EPSILON0 = 1/LIGHT_C/LIGHT_C/MU0; # Permittivity of free space
BOLTZMAN_C = 1.3805e-23 * J/K;  # Boltzman's constant
AVOGADRO_C = 6.023e23 *molecule/mole; # Avogadro's number of molecules
GRAVITY_C = 6.673e-11 * N*m*m/(kg*kg); # Newtons gravitational constant
GAS_C = BOLTZMAN_C*AVOGADRO_C;  # Gas constant
INFINITY=1.0e38;                # darn big number;
eCHARGE = 1.602e-19*C;          # Charge of an electron
EARTH_G = 9.80665 * m/(s*s);    # Earth's gravitational field, somewhere
eMASS = 9.1095e-31*kilogram;    # Electron rest mass, I suppose
pMASS = 1.67265e-27*kilogram;   # Proton mass

```

constant based  
conversions

eV = eCHARGE \* V;

keV = 1000\*eV;

MeV = 1e6\*eV;

GeV = 1e9\*eV;

TeV = 1e12\*eV;

PeV = 1e15\*eV;

EeV = 1e18\*eV;

lyr = LIGHT\_C \* yr; # Light-year

oersted = gauss/MU0;

subtly dimensionless  
measures

rad = radian;

srad = steradian;

deg = radian\*1.74532925199433e-2;

degrees = deg;

grad = 0.9\*deg;

arcmin = degrees/60.0;

arcsec = arcmin/60.0;

light quantities

cd = candela;

lm = candela\*steradian;

lumen = lm;

lx = lm/meter^2;

lux= lx;

miscellaneous rates

gpm = gallon/minute;

time variant  
conversions

MINIMUMWAGE = 4.75\*US/hr;

SPEEDLIMIT = 65\*mi/hr;

Conversions we'd like to see, but probably won't:

milliHelen = beauty/ship;



## Brief History of ASCEND

ASCEND is an acronym which stands for **Advanced System for Computations in ENgineering Design**<sup>1</sup>. The name ASCEND first appeared in print in 1978. The ASCEND programs are a series of modeling systems that Arthur Westerberg and his graduate students at Carnegie Mellon University have developed since that time.

### ASCEND I

Dean Benjamin developed the first ASCEND system. It was an interactive system in Fortran. Chemical engineering students at Carnegie Mellon University used this system from about 1978 to 1982 to carry out multicomponent flash calculations. It supported the senior design project.

### ASCEND II

Almost in parallel, Michael Locke developed the ASCEND II simulation system for his PhD thesis [1981]. ASCEND II allowed users to create models by configuring them using predefined types of parts. System maintainers defined the library of types, each in the form of seven handcrafted Fortran subroutines. These routines computed the space needed for the data when instancing a part, generated numerical values for the partial derivatives and the residuals of the equations that the part instance provided to the overall model, generated proper variable and equation scaling and the like. Michael Locke used this system to create models involving a few thousand equations to test variants of the Sequential Quadratic Programming algorithm. Tom Berna and he developed for optimizing structured engineering systems. Selahattin Kuru also used ASCEND II to generate and test solution algorithms for dynamic simulation that he subsequently developed for his PhD. Two companies used the software architectural design of ASCEND II to create their own internal equation-based modeling systems.

Experience at this time demonstrated that models involving several thousands of equations were solvable and could even be efficiently optimized. The question of interest moved from how to solve large equation-based models to how to aid an engineer to pose them, debug them and get them to solve.

In 1983 Dean Benjamin proposed the first version of a modeling language for posing complex models. Larry Gaydos and Art Westerberg further developed this language in the spring of 1984.

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1. ASCEND originally stood for “Advanced System for Chemical ENgineering Design” but the second generation system and following are not discipline specific, thus the name change.



## ASCEND III

In 1984 Peter Piela undertook a PhD project with Art Westerberg to “reduce the time needed to create and solve a complex model by one order of magnitude.” He developed what became ASCEND III. He had the help of two Carnegie Mellon University undergraduate students, Tom Epperly and Karl Westerberg, and of Roy McKelvey, a member of the faculty in the Design Department in Fine Arts. This team developed this system on the Apollo workstation and in Pascal. It comprised three parts: a modeling language and compiler, an interactive user interface and a suite of solvers. The language used object-oriented principles, with the exception of hiding of information. Modelers define types to create model definitions. A type (called a model in ASCEND) is a collection of variables and complex parts whose types are previously defined and the definition of the equations that model is to supply. A model can also be the refinement of a previously defined type. The language fully supported associative arrays and sets. For example, a distillation column is an array of trays. It also supported deferred binding by allowing one to reach inside a part and alter one of its parts to be a more refined type. The language and its compiler obviated the need to have a system programmer write the seven subroutines needed in ASCEND II.

The interactive user interface supplied the user with organized access to the many tools in the ASCEND III system. There were tools to load model definitions, to compile them, to browse them, to solve them, to probe them, to manipulate the display units (e.g., ft/s) for reporting variable values, to create reports and to run methods on them. One could even point at a part and ask that it be made into a more refined type (triggering the compiler to restart). As previously solved values were not overwritten, they became the starting point for the more complex model. Thus one could creep up on the solution by solving more and more complex versions of a model. Many of the tools were there specifically to aid the user in debugging their models as they tried to solve them. A tool could tell a user that the model appeared to be singular and why. Another set of tools aided in picking a consistent set of variables to fix before solving. Browsing allowed the user to look at all parts of the model. It was easy to check the configuration of a model. One could ask that parts of a model be solved one at a time.

Experience by Peter Piela, Oliver Smith, Neil Carlberg and Art Westerberg with ASCEND III demonstrated very clearly that *skilled* modelers could develop, debug and solve very complex models much more rapidly than they could with previously available tools, easily meeting the original target of a order of magnitude reduction in time required.

## ASCEND IIIc

In the fall of 1992, Kirk Abbott and Ben Allan approached Art Westerberg and said they wanted to convert the ASCEND III system from Pascal into C. They would also use Tcl/Tk for the interface. With these changes, the system would then run on most Unix workstations. Tom Epperly and Karl Westerberg had already created a C version for the compiler and solver. Abbott and Allan wanted to do this conversion even after they were warned that converting the system would take time that they could be using to do more apparently relevant work to complete their PhD theses. They insisted<sup>2</sup>. They were aided by Tom Epperly who, although located remotely, worked with them on the compiler. In eight months and putting in excessively long hours, they had a working system that could mimic most of the capabilities of the ASCEND III system.

Several students and a few people outside CMU could now use the system for modeling. Bob Huss and Boyd Safrit performed the hardest testing when they used ASCEND IIIc to model nonideal distillation processes. They developed and solved models involving up to 15,000 equations. Using a rudimentary capability for solving initial value problems, Safrit also solved dynamic models.

With use came the recognition of a need for improvements.

Attempts to teach ASCEND to others showed that it was a great system to speed up the modeling process for experts. Nonexperts found it nearly impossible to reuse models contained in the ASCEND libraries. The library for computing the thermodynamic properties of mixtures was particularly elegant but almost impossible to reuse. Modelers would reinvent their own properties models quickly, unable to use the library models.

Models larger than about 17,000 equations took more space than our largest workstation could provide. The models by Huss and Safrit were pushing the limits. Abbott and Allan established the goal to increase the size possible by a factor of at least ten, i.e., to about a quarter of a million equations. ASCEND needed to solve models more quickly. Without counting the increases coming from faster and larger hardware, the goal here too was a factor of ten. If solving were to be that fast, then compiling would stand out as unacceptably slow. The goal: ten times faster.

Abbott, with Allan, exposed a new style for modeling in ASCEND. He created prototypes of the various repeating types that occur in a model. The compiled equations and other data structures to define these prototypes then became available for all subsequent instances of parts that were of the same type as the prototype. Only the instance data

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2. It should be understood that Art Westerberg was thrilled they insisted on doing this conversion.

needed to be developed separately. Demonstrated impact on compile times was dramatic.

Abbott, with Allan, looked at how to speed up the solving times. The new twist was to use the model structure as defined by the model definition to expose a global reordering for the model equations before presenting them for solution. The time to solve the linear Newton equations as the inner loop of solving nonlinear equations dropped by factors of 5 to 10.

## ASCEND IV

Ben Allan has taken a lead role and worked with Mark Thomas, Vicente Rico-Ramirez and Ken Tyner to produce the next version of ASCEND, ASCEND IV. Playing the role of tester, an undergraduate, Jennifer Perry, demonstrated that Allan's introduction of parameterized types dramatically increased the reusability of the model libraries, converting it into an almost automatable exercise. Adding language constructs to permit the modeler to state what constitutes misuse of a model leads to the system generating diagnostics the model itself defines. Allan also completely revised the data structures and the interface between ASCEND and its solvers so that adding new solvers is much less work and so the solvers in ASCEND themselves become separable from ASCEND and usable by others.

Allan also defined the addition of NOTES to ASCEND which are like methods except they are not understood by the ASCEND system itself. Rather they can be passed to programs outside ASCEND. An example includes documentation notes which a documentation manager can use to compose answers to queries about what is in an ASCEND model. Another is a note that contains a bitmap description of a part that an external package could use to draw a symbol of that part.

ASCEND IV can now handle discrete variables and constants (logical, binary, symbolic, and integer). It supports the solver directing that parts of the model be excluded when solving such as when solving using implicit enumeration (dynamic model modification). CONOPT is now attached for optimization. The standard solver is rapidly becoming much more robust. ASCEND IV can generate a GAMS model corresponding the ASCEND model, giving access to solvers GAMS has that ASCEND does not.

While not quite there just yet, the goal to compile and solve 250,000 equations on a 150 megahertz workstation having about 250 megabytes of fast memory in a few tens of minutes is in sight.