Bil Reference Manual

The documentation for Bil 2.4 A modeling platform based on finite element/volume methods

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1 Introduction

Bil is a modeling platform based on finite element/volume methods dedicated to coupled problems involved in environmental engineering, geomechanical engineering, material sciences, etc.. Bil is distributed under the terms of the GNU General Public License (GnuGPL). The source code can be downloaded at

http://perso.lcpc.fr/dangla.patrick/bil

Bil is intended to be used by students, engineers or reseachers to work out problems or develop their own models. Bil is written in C language. Thus provided that a C compiler was installed, it can run on any OS. However the complete installation of Bil (including documentations) can be achieved on Linux-based OS only (Debian, Ubuntu, etc..).

Bil is developped for 1D, 2D and 3D problems. It doesn't include a mesh generator and postprocessing treatment of outputs. However it can read mesh files created by the open-source free software Gmsh downloadable at http://www.geuz.org/gmsh/. The output files created by Bil for 1D problems can be used easily by some plotting programs as Gnuplot. But as a general rule, post-processing options create output files which can be used by Gmsh for post-processing treatments (see Bil options).

2 Running Bil

Bil can be run non-interactively only. To compute the solution of a problem described in the input data file my_file, type:

bil [options] my_file

The input data file format is described in the following section. Without any option, running Bil will create the output files my_file.pi and my_file.ti (see Chapter 4 [Output files], page 9). If there is no input data file named my_file, Bil will open this file in writing mode and will ask the user for the input data. With the option -h (help) there is no input data file to give. The command

bil

(alone) displays the available options.

The way Bil will run may depend whether there are some other files (see Chapter 5 [Other files], page 11). The most important of them, namely my_file.graph.iperm, defines the way nodes are renumbered so as to optimize the resolution algorithm. Except in 1D problems, it is very important, if not essential, to create this file with an aquedate option, e.g. -iperm, (see Bil options).

3 Input data file format

The file my_file, as mentionned above, provides the input data of the problem to be worked out. A list of 4 capital letter key-words organizes the inputs in several groups such as mesh, material properties, boundary conditions, etc. For example the key-word GEOM is followed by the inputs defining the dimension and the symmetry of the problem. The full list of key-words which must appear in my_file is given in the table below. Any line begining with **#** is considered as comments and skipped.

key-word	description	
GEOM MESH MATE	dimension and symmetry of the problem mesh material properties of the material index 1	
MATE	material properties of the material index n	
FLDS	fields i.e. space functions	
INIT	initial conditions	
FUNC	time functions	
COND	boundary conditions	
LOAD	loads	
POIN	define some points for output files	
DATE	define the dates for output files	
OBJE	objective variations of the main unknowns	
ITER	convergence criteria of iterative process	
TIME	time steps calculation	

An on-line help is provided by typing:

bil -h

4 Output files format

Each run produces 2 sets of output files. In the first set, output files are named:

my_file.pi

where i is an integer ranging from 1 to the number of points defined by the key-word POIN. There is no file if there is no points. These files provide the results obtained at the specified points. The first column contains the times at which the results have been obtained. The other columns contain the value of specific quantities as implemented in the model defined in the key-word MATE.

In the second set, output files are named:

my_file.ti

where i is an integer ranging from 0 to the number of dates defined by the key-word DATE. These files provide the results obtained at the specified dates. The three first columns contain the three coordinates of nodes. The other column contain the value of the same quantities as those contained in the first set of files.

Some lines of these 2 sets of files are commented as indicated by the character **#** in the first column. These comments provide some informations about the nature of the computed quantities found in the following lines.

5 Other files

Bil produces some files and sometimes can read and use some other files. The name of these files are formed with the name of the input data file and suffixes, my_file.suf, in the same way as the output files. They are listed in the table below.

fichier	description
my_file.t <i>i</i>	output files related to date index i
my_file.pi	output files related to point index i
my_file.posi	view i to be read by Gmsh
my_file.msh	Gmsh mesh file
my_file.graph	mesh graph
my_file.graph.iperm	inverse permutations file
my_file.sto	storage file
my_file.cont	continuation file (see below for explanations)
my_file.conti	continuation file (see below for explanations)

The files my_file.cont and my_file.conti allow to continue a previous computation or to resume an interrupted calculation achieved with a previous input data file (my_ previous_file). With my_file.cont the process doesn't go through the initialization stage (ComputeInitialState, see below) so that the calculation continues as if there hadn't been interruption. With my_file.conti the process goes through the intitialization stage so that some variables of the model can be re-initialized (e.g. strain variables can be reset to zero). To do so, copy the file my_previous_file.sto in my_file.cont (or my_file.conti) and run bil with my_file as a new input data file in which you will have defined some additional dates beyond the last date defined in my_previous_file.

6 Models

Mostly the concept of model refers to the constitutive equations or complementary laws that are needed to mathematically end up with a well-posed problem. However we need more informations, here, regarding the numerical methods that are used to handle complex problems and geometries. These methods will be implemented in a single file whose basename will identify the code name of the model. The concept of model will then be extended to fit with the set of informations pertaining to:

- The number and kind of equations to be solved
- The complementary laws that are needed to have a well-posed problem
- The informations associated to the numerical methods employed

Models aim at addressing the behavour of the material at the scale of one finite element. Therefore the methods defined in object "Model_t" aim at computing matrix, residual forces, outputs and so on, for the nodes of one element. The object "Element_t" is therefore the main input entry of all methods of object "Model_t".

A short description of the available models can be displayed by typing:

bil -m

7 How to develop a new model ?

To tell Bil to account for a new model, you just need to create a new file, e.g. my_model.c, in the folder ModelFiles and add the basename of this file, namely my_model, to the list of the available models found in ListOfModels.inc. This model will be taken into account automatically the next time binary files will be created.

To help you in creating this new file, it is recommended to learn from already existing files. This file should contain at least the 11 methods of the Model class-like structure. These methods are listed in the table below.

method

description

SetModelProp ReadMatProp PrintModelProp DefineElementProp ComputeInitialState ComputeExplicitTerms ComputeMatrix ComputeResidu ComputeLoads ComputeImplicitTerms ComputeOutputs Set the model properties Read the material properties Print the model properties Define some properties of the element. Compute the initial state Compute the explicit terms Compute the matrix Compute the residu Compute the loads Compute the implicit terms Compute the outputs

8 Examples

8.1 Drainage of a column

This problem is governed by the Richards' equation. A 1 meter high sand column is initially satured. The liquid pressure is initialized as: $p_l = p_{atm} - g(x - 1)$. At t = 0 we drained the column from the bottom by imposing the pressure to $p_l = p_{atm}$. The input data file is given below.

inputs	comments	
# Drainage of # a sand column	You can write some comments by beginning any line with $\#$.	
GEOM 1 Plan	Geometry of the problem 1D pb, plane symmetry	
MESH col.msh	Mesh The mesh is read in this file (format Gmsh). This mesh consists in a 20 elements mesh between 0 and 1. There are 2 regions. The region 1 is the point at 0. The region 2 is the line between 0 and 1. There is 1 material.	
MATE Model = m1 gravite = -9.81 phi = 0.3 rho_l = 1000 k_int = 4.4e-13 mu_l = 0.001 p_g = 100000 Curves = tab	Material 1 code name of the model gravity porosity fluid mass density intrinsic permeability fluid viscosity gas pressure in the file tab, there 3 columns: $p_c S_l k_{rl}$	
FLDS 2 Type = affine Value = 1.e5 Gradient = -9.81 Point = 1.	Fields 2 fields affine field defined by $10^5 - 9.81 * (x - 1)$	
Type = affine Value = 1.e5 Gradient = 0. Point = 0. INIT 1 Region = 2 Unknown = p_1 Field = 1	constant field equal to 10^5 Initial conditions 1 initial condition in the region 2, $p_l = 10^5 - 9.81 * (x - 1.).$	
FUNC O	Time functions $f(t)$ here there is no function	
COND 1 Region = 1 Unknown = p_1 Field = 2 Function = 0	Boundary conditions 1 boundary condition in the region 1, $p_l = f(t) * 10^5$ (by default $f(t) = 1$)	

LOAD	Loads
O	there is no load
POIN	Points where we want outputs
O	no points
DATE	Dates where we want outputs
2	2 dates
0. 1800000	$t_0 = 0$ and $t_1 = 1800000$
OBJE p_l = 1000	Objective variations of unknowns objective variation $\Delta p_l = 1000$
ITER Iterations = 20 Tolerance = 1e-10 Repetitions = 0	Parameters for the iterative process 20 iterations the tolerance is 10^{-10} no repetition
TIME	Parameters for time steps calculation
Dtini = 1	initial time step equal to 1.
Dtmax = 3600	maximum time step equal to 3600.

9 Versions

News in 2.3: Use of C++ compiler. Programmation is now extended to C++ langage. Hence the use of .cpp and .hpp files is allowed. A new object "Exception.h" has been created to handle exception mechanisms such as interruption, floating point error. In such occuring event the program handles the event, saves outputs and exits in a clean way. Introduction of the curve builder "Expressions", based on the evaluation of mathematical expressions obtained from AnaGram (www.parsifalsoft.com). In this version a shared library, libbil.so, from the genuine sources of Bil is created and installed in the machine. Linkage with other possible external libraries is possible. A new extension of input file ".conti" has been introduced so that we can load the solution from a previous calculation and continue this calculation while going through the initialization stage (see documention).

News in 2.2: WARNING, the outputs of FVM_ComputeIsotropicConductionMatrix() and FVM_ComputeMassAndIsotropicConductionMatrix() have been modified see examples in models using these methods. New objects have been introduced: Views, TextFile, CurvesFile, MatrixStorageFormat. Introduction of the curve builder "Evaluate", based on the evaluation of mathematical expressions obtained from Snippets (www.brokersys.com/snippets/). Some bugs fixed, e.g. those in "Buffer_FreeFrom". Extended field delimiters in curves files. Some new models.

News in 2.1: A class-like structure Buffer_t has been introduced as a circular buffer. It can be used in any functions to compute vectors or tables of any type. Numerical Methods, such as FEM or FVM, have been implemented in files such as FEM.[c,h] and FVM.[c,h]. These methods are viewed as object with class-like structures FEM_t and FVM_t. In these files methods have been implemented such as FEM_ComputeMassMatrix and can be used directly in models provided that header has been included. The folders "Common" and "Main" have been created. We moved the main files in "Main" and files of common use, like "Buffer.c", in "Common". DataBases have been created which can be used in model files. The matrix storage format have been implemented in a separate file "MatrixStorageFormat.h".

News in 2.0: The code has been rebuilt to an object-oriented programming code. However the code is still implemented in C language. Class-like structures have been introduced and implemented in separated files. Each class-like structure has attributes and pointers to functions (see e.g. structure Model_t). Moreover some improvements have been introduced. The file names of models (in the folder "ModelFiles") can be chosen abitrarily. The class-like structures "NodeSol_t" and "ElementSol_t" are two linked lists which contained the nodal and elemental solutions. A new pointer to "double" type in "ElementSol_t" can be used to store constant terms.

Nouveautes dans 1.8: creation d'une structure "modl_t" contenant les methodes (taches elementaires) i.e. des pointeurs sur fonctions.

Nouveautes dans 1.7: option "base" ajoute dans le Makefile. Qqs nouveaux modeles. Option "Relative" possible dans OBJE, donnee apres la valeur. Chargement automatique des modeles, il n'est plus besoin de modifier le fichier "xmod.c". Les fonctions "dm,qm,tb,ch,in,ex,ct,mx,rs,so" sont renommees "dmNB,qmNB,tbNB,chNB,inNB,exNB,ctNB,mxNB,rsNB,soNB". Les titres des modeles ainsi que des exemples de donnees sont geres par les fonctions "qm1, qm2 ...". Options nouvelles de la ligne de commande. "bil -m" affiche les titres des modeles. "bil -m I" affiche un exemple de donnees du modele I. Possiblite d'avoir des elements d'ordres multiples (1 pour chaque inconnue). Cette possibilite est geree par les tableaux "el.pin" et "el.peq". Une valeur negative associee a un noeud et une inconnue n'est pas prise en compte comme inconnue globale. Possibilite de creer de nouvelles fonctions d'interpolation au niveau des modeles dans "tb1, tb2, ... " avec la fonction "creer_interpolation(...)". Possibilite de definir des champs aux points d'intersection d'une grille dans l'espace avec l'option "Type = grille" dans CHMP (l'option Type = affine restant par defaut).

Nouveautes dans 1.6: bug corrige dans "sauvep". Les fichiers "mod.c" et "mod.h" sont renommes "xmod.c" et "xmod.h". Le specificateur de type "void" est remplace par "int" pour les fonctions "ex1(), ex2() ..." definies dans "m1.c, m2.c ..." et pour l'identificateur de type "ex_t" defini dans "defs.h". On utilise ce retour dans les fonctions "explicite()" et "algorithme()" definies dans "calc.c". Compatibilite avec le format de maillage version 2.0 de GMSH.

Nouveautes dans 1.5: definition des types de fonctions "dm_t, "qm_t", "tb_t", etc... Reorganisation du fichier "mod.c" par l'introduction de tableaux de pointeurs de fonctions pour simplifier l'introduction de nouveaux modeles. Possibilite de charger la librairie de SuperLU (voir le fichier "make.inc"). Creation de l'option de la ligne de commande "-m slu" qui permet d'utiliser la methode de resolution proposee par SuperLU a condition d'avoir construit, auparavant, le fichier des permutations inverses a l'aide d'un programme adapte comme Metis.

Nouveautes dans 1.4: reorganisation et creations des fichiers : calc.c, lecdo.c, postt.c, renum.c. Suppression des variables statiques. Creation d'une structure de donnees "dnns_t", d'une structure pour la matrice "mtrx_t", d'une structure pour la solution "sltn_t", d'une structure pour la renumerotation "nume_t". Amelioration de certaines fonctions de lib.c. Suppresion de rssurf (pris en charge par rsmass).

Nouveautes dans 1.3: nouveaux modeles inclus. Qqs ameliorations apportees dans le calcul de dt (fonction pasdt). Apport de nouvelles info dans les structures mate_t, elem_t: en particulier neq,eqn,inc dans mate_t. Creation de la structure node_t et suppression du pointeur no_x. Cette version permet de prendre en compte des modeles reposant sur un nombre d'equations et sur des natures d'inconnues pouvant varier d'une region a l'autre. Cette version realise la continuite des equations en fonction de nouvelles informations contenues dans les modeles comme le nombre d'equations et les noms (predefinis) de chaque equation et inconnue associee. En consequence la structure du fichier a ete (legerement) modifiee pour prendre en compte ces informations, notamment dans les mots-cles INIT,COND,CHAR,ALGO (voir l'aide en ligne). Pour plus de clarte les donnees relatives aux variations objectives des parametres ont ete deplacees du mot-cle ALGO dans le nouveau mot-cle OBJE.

Nouveautes dans 1.2: creation d'un repertoire exemples. Simplication de la gestion des elements de surface (suppression de SURF). Creation de champs (mot-cles CHMP). Gestion des interruptions dans le calcul de la matrice.

Nouveautes dans 1.1: creation des pages info et de la doc sous differents formats (ps,pdf,txt). Creation des repertoires bin et lib. Amelioration de la procedure d'installation. Simplification du jeu de donnees par la definition des regions de maillage.

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Version 2, June 1991

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