Response to editorial and reviewer Comments for “Simulating Coupled Surface-Subsurface Flows with ParFlow v3.5.0: Capabilities, applications, and ongoing development of an open-source, massively parallel, integrated hydrologic model”

We would like to thank the Executive Editor, the Handling Topical Editor, and the referees for the comments and suggestions intended to improve our manuscript. The minor comments provided by the referees were particularly helpful in bettering specific sections of the manuscript, including some word omissions, typographical errors and clarifications, which can be difficult to spot in one’s own manuscript. The revised manuscript addresses the concerns raised by the referees and we are confident the Editor will find it suitable for publication without an additional external review.

Please find below our revision, which includes detailed point-by-point response to all referee comments. The response to the referee comments are structured as follows: (1) comments from referees, (2) author's response, and (3) author's changes in a marked-up manuscript version showing all the changes made. We are glad to discuss any aspect of the revision with you, the handling topical editor, or any of the referees if questions arise. The original text of the comments is below in plain face font and our responses are given in bold face italic font for maximum clarity. We have included references to specific line numbers regarding all changes and these line numbers refer to the “tracked changes” version to make it as simple as possible to evaluate the changes in the context of the previous submission.

The Executive Editor’s Comments:
This is an executive editor comment highlighting the ways in which this manuscript is not currently compliant with GMD policy on code and data availability. In this case, there is just a single technical issue which needs to be remedied in the revised submission:
1. Github URLs. Github is an excellent development platform, but it lacks the features required of an archive. GitHub themselves tell authors to use Zenodo for this purpose. The authors should follow the procedure detailed there to archive the exact version of the software used to create the results presented: https://guides.github.com/activities/citable-code/. The resulting Zenodo repositories present the correct bibliography entries to use. Further details on code and data availability requirements are in the GMD model code and data policy: https://www.geoscientific-model-development.net/about/code_and_data_policy.html. The reasons for the policy and more detail are provided in this editorial: https://doi.org/10.5194/gmd-12-2215-2019.

Authors Response
We greatly appreciate the Editor for his comments. The Editor’s directions that we adhere to GMD’s policy on code and data availability and use Zenodo has been considered. We have made modifications to that effect in the revised version of the manuscript.
See Lines 985-987 in the revised manuscript.

Anonymous Referee #1 Comments
Authors have summarized major advances in development of an integrated hydrologic
atmospheric model (ParFlow.*) for simulating terrestrial hydrologic processes. The paper is a nice summary of authors’ effort in the past 3 decades on ParFlow development, and its coupling with land surface, atmospheric and reactive transport models. As authors state, the information presented here has been previously published as part of software manuals and papers published by the developers. Therefore, this manuscript provides a valuable resource for the users to learn about the model functionality.

However, it would be more useful if authors consider adding the following information:

1. The paper falls short in describing capabilities of ParFlow in comparison to other integrated hydrologic models such as CATHY, HydroGeosphere, etc. This will help users with model selection for a particular application.

Authors Response

We agree with the reviewer that it is important to understand how one integrated model compares to others but note that this has been done recently. The integrated model inter-comparison project was a series of workshops where the developers of many of the contemporary integrated models got together and designed test cases for comparison. Some of these results are highlighted in Maxwell et al. 2014, Kollet et al. 2017 and Koch et al. 2016. Given the extensive work presented in these existing papers, and the extraordinary effort it takes to run such simulations, we feel that including a detailed comparison here is beyond the scope of the review, but we have emphasized these inter-comparison resources for the readers.

2. It would be very useful if authors could describe future model development. What is next?

Authors Response

Absolutely, and we are glad to add these details. We had mentioned the availability of a software development and sustainability plan to improve the capabilities of ParFlow in the last paragraph of the manuscript, but details on future directions were an unintended omission in the draft. We have added text concerning the code developers next plan in improving the code such as new formulations of both kinematic and diffusive wave approximations, and advanced parallelization support (GPU’S and heterogeneous compute architectures) (Lines 968-970). We also note that the development branches and forks can be viewed on the GitHub page (https://github.com/parflow).

3. Despite improved parallel efficiency for large scale application, model application for large domains is computationally intensive. Can authors provide further guidelines for model set-up (table of inputs), initialization and calibration? Are there any efforts underway to improve computational time?

Authors Response

We completely agree with the sentiment of this comment; 3d Richard’s equation and overland flow is computationally expensive and even the best parallelism cannot change that...
fact. One of the challenges to providing guidance here is that ultimately it is up to the user to determine what level of complexity is appropriate for their problems and what portions of the problem to simplify. That level of complexity combined with the heterogeneity of specific elements dictates how long it takes the model to iterate, which is the main factor during initialization and also for calibration. The long run times often inhibit or prevent calibration efforts, so many integrated models to date have not been calibrated and instead are used as virtual hypothesis testing platforms; we’re not saying this is necessarily a “best practice in modeling” (we think sensitivity analysis and uncertainty quantification should always be included in planning models) but uncalibrated models that are based directly on the physics can still provide useful results about process interactions. As for initialization, the common practice is to “spin-up” (e.g. Seck et al. 2015) the model to a steady- or pseudo-steady (in the case of a transient forcing) state, but even for one specific kind of model there are multiple workflows that can be used, so this could not be one table rather a tangled spider web. We absolutely agree that guidance is useful in these areas but respectfully disagree that an overview of a particular model is the best place for such guidance. However, recognizing the importance of this comment, we have added clarifying text noting that the studies involving ParFlow outlined in Table 1 provide a wealth of knowledge regarding domain setup (Lines 536-543). Since these are all specific applications, their information will likely be very useful to modelers trying to build a new domain for a similar application as they are setting up and planning their model. As for efforts to improve efficiency, the numerical solvers developed for ParFlow now comprise the core of the Hypre and Sundials solver libraries (the common PFMG preconditioner in Hypre stands for ParFlow Multi-Grid) and are currently the most numerically efficient solvers available. As such, many ParFlow developers are currently focused on increasing performance on heterogenous compute architectures, as noted above in our response to comment 2.

4. While authors summarized various application of the model in Table 2, it would be great if they can present a simple case study that compares computational time as different components are added from land surface to the atmosphere, and show how simulated outputs have been improved compared to observations.

Authors Response
We are in complete agreement that this is common question and that it would be great to have such a table, but in practice it is extremely difficult to provide any general examples like this because of the nonlinearities in the different parts of the solution of the system. For example, the geochemical systems in ParCrunchFlow compared to the nonlinear snowmelt processes in CLM, or the switching on or off of the overland flow routing. Each of these has a different cost, and when coupled to other portions of the problem the numerical cost (i.e. solver time) of their interactions is, for all intents and purposes, impossible to generalize. The best one could do is to highlight a few specific, contrived examples that could never be considered generalizable. The best estimate of how solver time changes would be considering the number of floating-point operations per iteration, but even this cannot be predicted
because the nonlinear system evolves over time so the number of iterations for convergence constantly changes. The relevance of this comment is without question, but it is also deceptively difficult to resolve. Many of the studies presented in Table 1 include computational times for problems with different complexities where ParFlow was used, and this may be able to help readers infer how runtime will change; we have added comments to this affect (Lines 518-527). If one could predict how adding processes would affect runtime that would be an incredible advance for scientific research, but we are aware of no such advances. The best we can offer the reader is a brief summary of these thoughts and their justification (Lines 518-527), which we think the reviewer will agree will be a useful discussion for the readers

Minor Comments

Lines 82-85- The differences between the integrated approach and indirect approach is not clear. Please explain.

Authors Response
This is a good point. After reviewing the original text, we completely agree that this was not clear, and have modified to accommodate suggestions. For example, further descriptions on how ParFlow employ these approaches pertaining simulation of flows in surface and subsurface domains (Lines 87-91) of the revised manuscript.

Line 94 – Kollet et al. (2010) does not seem to be a suitable reference here as the focus of the paper is on parallel efficiency. Please refer to Kollet and Maxwell (2008), Water Resources Research instead.

Authors Response
We absolutely agree; clearly, the citations were mixed up. This has been corrected (Line 100 in the revised manuscript) and we’re grateful for the reviewer’s sharp eyes.

Line 139 – Is the variable vertical discretization only possible with the terrain following grid option in ParFlow?

Authors Response
Variable vertical discretization can be used with any domain/grid; however, it usually makes the most sense to do so with a terrain following grid since this is commonly used to increase the resolution of the shallow soil layers (Lines 150-151 in the revised manuscript).

Line 155 – Remove “of” from “relative of saturation”
Correction made See Line 168 in the revised manuscript.

Line 171- According to equation 4, units of Darcy flux should be LT-1.
Authors Response
A rather embarrassing mistake on our part that has been corrected. See Line 170 in the revised manuscript.

Section 2.3. Add information regarding flow routing approach. For example, does the new version support D8 flow direction?

Authors Response
D4 flow direction is implemented in ParFlow. Notes have been added to explain this (Lines 220-222).

Line 194 – Move “slope” before the “(gravity forcing term)”

Authors Response
Change made. See Line 211 in the revised manuscript.

Line 254 – Add “relative” to Si

Authors Response
“Relative” has been added. See Line 275 in the revised manuscript.

Line 742- To main consistency, write units.

Authors Response
The units are now provided for all variables presented. See Lines 798-800 in the revised manuscript.

Anonymous Referee #2 Comments
The motivation (and objective) of the paper is stated as (Lines 71-74) : “The purpose of this manuscript is to provide a current review of the functions, capabilities, and ongoing development of one of the open-source integrated models, ParFlow, in a format that is more accessible to a broad audience than a user manual or articles detailing specific applications of the model”. I am very familiar with integrated hydrologic models, but not with ParFlow, and I therefore belong to the target audience. However, after very carefully reading the paper (some sections more than once because they could be clearer), I conclude that the paper does not reach its objective and does not provide a clear review of the code’s functions, capabilities, and ongoing development. Overall, the organization and writing should be improved to make the text much clearer. Some sections provide too much information on peripheral details and too little on some important points. That is especially the case for the coupling section (section 5), which does not provide a clear picture of the code’s capabilities with respect to its coupling with other codes. I provide below more detailed comments on specific sections of the paper.

Title
The title is not representative of the content of the paper.
• With respect to coupling, a good portion of the paper focuses on describing the coupling of ParFlow with other codes. The focus is therefore not so much on coupled surface and subsurface flow as the title suggests. Actually, the surface and subsurface coupling could be described more clearly (see comment below).
• The capabilities are described but the paper does not provide a clear picture of the applicability limits of the model.
• The ongoing development is not really addressed. The paper rather lists past developments

Authors Response
The reviewer’s statement that “The title is not representative of the content of the paper” is to some extent a reasonable criticism and appears to be constructive in nature. However, we are faced with a choice and must decide amongst many possible tradeoffs. While it has been coupled to many different codes, the main function of ParFlow is to simulate coupled overland (surface) and subsurface flows, which is reflected by the title. We concede that a different title might be able to capture the complex nature of all the possible simulations but stop short of changing to a complete list of all the couplings included. However, to address these concerns, we have clarified the content and scope in the abstract (Lines 35-38).

We think the applicability of ParFlow has been well discussed in section 6 (Discussion and Summary) highlighting challenging hydrologic projects or research works where ParFlow was used. For example, the code has been applied to simulate surface and subsurface flows at varying spatial scales i.e. from mouth of continental river basins at high resolutions, evaluate relationship between topography and groundwater flow, assess resilience of water resources and anthropogenic stressors, and simulate atmospheric, surface and subsurface energy and water budgets in coupling with other models. The impetus is on the individual modeler to select a model that is applicable to their task, and we think the paper presents sufficient example for readers to inform their own opinions on when the approach is applicable.

To the final point, we completely agree that we should have included these details before and thank the reviewer for catching this omission. The manuscript has been revised to add ongoing development plan in the last paragraph such as incorporation of new formulations of both kinematic and diffusive wave approximations, and advanced parallelization support (GPU’S and heterogeneous compute architectures). Lines 968-970 in the revised manuscript.

Introduction
The introduction does not fit with the purpose of the paper, which is to present an overview of ParFlow’s capabilities.
• There are some very broad statements on integrated hydrologic models (IHMs) in the first paragraph that are not really required since the intended audience will likely be already aware of IHMs and will not need to be convinced of their usefulness.

Authors Response
We thank reviewer for this critical evaluation, and think we understand their intent, but we disagree that the general statements on integrated hydrologic models (IHMs) are superfluous. Our intended audience is not only those well-vested or with broad knowledge in numerical or hydrologic modeling but include those looking to learn more about IHMs (e.g. graduate students) of whom we hope will consider ParFlow. We believe it would be useful to such potential code users and readers to know what IHMs are in broad strokes, what they do and why ParFlow belongs to that class of simulation platforms.
The second paragraph (lines 75-94) provides a short summary of ParFlow’s surface and subsurface flow capabilities. It is somewhat confusing to provide this summary in the introduction since the main goal of the paper is to provide a much broader overview of the code.

Authors Response
We think this is a good point. The intent was not to summarize coupled surface-subsurface flow simulation in the introduction section but provide introductory statements of such capability of ParFlow and then explain further in section 2 of the paper, which we did in sections 2.1, 2.2, and 2.3. Further, a brief introduction needed to be provided to lead our readers to what is to be discussed in the paper in terms of surface-subsurface flow simulation by ParFlow.

Lines 95-103 provide a list of previous studies but the description of the scale of application is confusing (large domains, small catchments, complex terrain, large watersheds, continental scale...). Also, the main conclusions or results of these studies are not mentioned. Just citing papers is not helpful. It would be better to comment on these studies to provide the reader with a clearer understanding of the code’s applicability. There are several other instances where a list of ParFlow applications is given, without much detail, (example are lines 132-139, lines 161-163, lines 870-875), which generates repetition.

Authors Response
It is completely true that scale can be arbitrary; this is an appropriate criticism and we concur with the reviewer. The manuscript has been revised according to the reviewer suggestion to give numerical evidences to the use of description of the scale of application such as large domains, small catchments, and large watersheds. This is fully exemplified in Table 2 where all of these descriptions are given values of lateral and vertical extents based on the referenced studies (Lines 103-107 and 111-113 in the revised manuscript).

Section 1.1 on the development history is interesting and relevant (although lines 132-139 can be removed).

Authors Response
We thank the reviewer for this suggestion, however, we’d be glad if this is reconsidered, because Section 1.1 gives a general trajectory of the code’s development based on periodic modifications and applications that assesses the code’s capabilities. Lines 132-139 (in the original manuscript) end the section with brief recount of some of the recent tested additional modifications and applications which were discussed in subsequent sections of the manuscript. So, we view the presentation in those lines very necessary in the manuscript.

Core functionality
It is not clear why variably-saturated and steady-state saturated modes are identified separately. Equation 1 is the transient variably-saturated flow equation and equation 3 is derived from the same equation by setting the time derivative to zero and both relative permeability and saturation 1.0. Why treat them separately, especially since a common solution method is used (line 148)? I would only present equation 1 to avoid confusion.

Authors Response
The two equations (1 and 3) were presented to elaborate the fact that the steady-state saturated flow can be derived from the variably saturated flow. This was done for the purpose of simplification and clarity of the equations, but we agree that additional clarification is needed here. However, ParFlow does include a direct solution option for the steady-state saturated problem that is distinct from the transient solver and this note has been added (Lines 188-191).
• Lines 179-185 are out of place and probably not necessary. If they are kept, they should go into an introduction. Same comment for lines 293-300.

Authors Response

The reviewer finds lines 179-185 and 293-300 (in the original manuscript) unnecessary and out of place, but we wish he takes a second look into that. We found it highly essential to begin each section or subsection with a brief introduction or background to lead our readers into what is it we’d be discussing in the said section of the manuscript. Moreover, large portion of the information provided in those lines (in the original manuscript) have been presented differently to some extent in the introduction section of the manuscript (Lines 51-59 revised manuscript). Lines 196-204 only elaborate notes introduced in section 1 of the manuscript.

• The description of the coupling between surface and subsurface (pages 8-9) is confusing and should be clarified. I think that there is two-way coupling in ParFlow but the text suggests that there is only flow from surface to subsurface (see lines 204-206: “To account for vertical flow (into the subsurface from the surface), a formulation that couples the system of equations through a boundary condition at the land surface becomes necessary”). Figure 1 suggests the same one-way flow direction.

Authors Response

This was poorly phrased on our part. There is a two-way coupling of the surface and subsurface flows in ParFlow and we have revised to make that clear (Lines 225; and 233-234).

• It is also not clear if surface and subsurface are coupled everywhere during a simulation or only at limited locations. Section 3.4, which describes the solution for the coupled surface and subsurface flow system, seems to suggest that surface flow is not solved everywhere (although I am not entirely sure because section 3.4 would have to be written more clearly).

Authors Response

Surface-subsurface coupling can occur anywhere in the domain during a simulation and it can change dynamically during the simulation. Coupling between subsurface and surface or overland flow in ParFlow is activated by specifying an overland boundary condition at the top surface of the computational domain, but this mode of coupling allows for activation and deactivation of the overland boundary condition during simulations where ponding or drying occur. Overland flow may occur by the Dunne or Horton mechanism depending on local dynamics. Overland flow routing (kinematic or diffusive wave) is enabled when the subsurface cells are fully saturated. Clarifications of these points have been made on Lines 464-471 in the revised manuscript.

• I do not see the usefulness of section 2.4. There is no evidence that the multiphase flow capabilities are used and the explicit time-weighting scheme used for transport is extremely restrictive for real applications, as well as the absence of dispersion or diffusion. It seems like these options are seldom used.

Authors Response

We agree with the reviewer to some extent, that the multiphase flow capabilities of ParFlow have had limited applications in recent times. However, we provided this functionality of the code to prompt or alert potential code users of the existence of that verified and working capability of ParFlow.

Equation discretization and solvers
• The writing style is clearer for this section, compared to the rest of the paper, but there are still some inconsistencies. For example, the method used to solve the variably-saturated flow equation is mentioned in 3 different places, but it is not consistent
• Lines 365-367: for variably saturated subsurface flow, ParFlow does this with the inexact Newton-Krylov method implemented in the KINSOL package
• Lines 372-373: For variably saturated subsurface flow, ParFlow uses the GMRES Krylov method
• Lines 409-410: For variably saturated subsurface flow, ParFlow uses the Newton-Krylov method coupled with a multigrid preconditioner

Authors Response
The different solvers mentioned in Lines 386-387 are all existing options in ParFlow capable of solving variably saturated subsurface flow equation. Inexact Newton is consistent across all but the preconditioners can be manually changed. The choice of a solver depends on the specific problem(s) being solved, and the code user may select which solver to use. For example, for a particular problem, one solver may provide faster convergence compared to the other. In that case, that solver may be the choice of the code user.

• Similarly, for saturated flow, it is written
• Lines 415-416: For saturated flow, ParFlow uses the conjugate gradient method also coupled with a multigrid method
• Lines 430-431: ParFlow uses the multigrid-preconditioned conjugate gradient (CG) solver to solve the groundwater equations under steady-state, and fully saturated flow conditions
• Either the conjugate gradient method coupled with a multigrid method and the multigrid-preconditioned conjugate gradient represent the same solution method (in which case there is unnecessary repetition) or they are different solution methods (in which case some more information is required).

Authors Response
Conjugate gradient method coupled with multigrid method and the multigrid-preconditioned conjugate gradient for saturated flow are different ways of presenting the solver in the performance of its function and not necessarily repetition.

Coupling
Section 5 on coupling is the section that requires the most careful revision.
• PF.CLM: It is mentioned that a modified version of CLM was incorporated into ParFlow. There is no clear description of the modified CLM (only some examples of capabilities, as listed starting on line 552). There is also no mention of the differences between the modified CLM and the original CLM published by Dai et al. (2003). Considering the aim of the paper, it would be useful to at least list the main capabilities and types of applications, instead of referring to previous work (lines 566-567). There is a mention of comparison to uncoupled models (line 588) but no identification of what the uncoupled models are. Also, since the modified CLM has been integrated into ParFlow, PF.CLM is not really a coupled model in the same sense as the other coupled models presented in section 5.

Authors Response
The difference in module structure between the original CLM and the modified version integrated into ParFlow was provided in the original manuscript. See Lines 620-631 of the revised manuscript.

It may not be feasible to enlist all the capabilities and application of ParFlow presented in the previous research works in details. We believe highlighting essential capabilities and/or applications such as the capability of PF.CLM to predict accurately root-depth soil moisture and referring readers and
potential code users to those resources where tested and specific applications are provided in detail would be useful.

The phrase “uncoupled model” simply meant a stand-alone model used in a simulation e.g. performing a simulation with CLM (land surface model) to compute soil moisture content without coupling with other model (ParFlow), then CLM is an uncoupled model in that regard. We have explained this in the revised version of the manuscript in Line 643.

- ParFlowE.CLM: The section mentions that a 3D heat transport equation has been added to ParFlow, which becomes ParFlowE. Since heat transport appears to be a core feature, why is it only mentioned here instead of being presented much earlier in Section 2? Is it because ParFlowE is a different ParFlow? Also, it is really not clear if the CLM used in ParFlowE.CLM is the same as in PF.CLM. Is ParFlowE available to use with the other models listed in Section 5?

Authors Response
ParFlowE was not included to section 2 because it is a modification made to ParFlow for a specific application. It is included in section 5 because ParFlowE is essentially ParFlow with 3D heat transport formulation addition and coupled to the CLM. It was explicit in the manuscript the original CLM by Dai et al., 2003 was used in coupling ParFlowE.

- ParCrunchFlow: That section is confusing. There is a description of CrunchFlow and its solution methods (lines 769-794) but it looks like only the reaction terms computed by CrunchFlow are used by ParFlow and the advective-dispersive transport capabilities are not used. If that’s the case, I would not describe all the CrunchFlow features, only those used. It would also be interesting to indicate why CrunchFlow’s advective-dispersive transport capabilities are not used and the advection-only capability of ParFlow is used instead, with its restrictive explicit time-weighting scheme. I assume that it’s a question of dimensionality but it is not clearly stated. Also, the reader has to guess that ParCrunchFlow is only applicable for subsurface simulations (it should be clearly stated). The whole section would need to be rewritten more clearly.

Authors Response
The document has been revised to highlight why CrunchFlow’s advective-dispersive transport capabilities are not used in the coupled model (ParCrunchFlow). ParCrunchFlow makes use of multidimensional advection capability of ParFlow instead of CrunchFlow’s advective-dispersive transport capabilities (up to two-dimensional). See Lines 861-863 in the revised manuscript. It has been made in the revised manuscript that ParCrunchFlow is applicable only in the subsurface. See Line 852 in the revised manuscript.

- The terminology used to describe the coupling of ParFlow with other codes is not consistent and can be confusing. There is a mention of offline and online couplings in section 5, which are fairly clearly described, but those terms are not used after that. It would be clearer if a constant terminology was used to describe the type of coupling.

Authors Response
The entire section 5 has been revised for consistency in the use of terminologies “online” and “offline” couplings. See Lines 635, 689, 736, 807, and 859.

Discussion and Summary
That section does not contribute much to the paper. Some sentences and statements are too general. One example is the first paragraph of the section.
The very last paragraph provides some practical information about ParFlow. From the point of view of a potential user or developer, it would be interesting to develop that aspect. For example, there is a mention that a software development and sustainability plan exists. It would be very interesting to provide a summary of that plan. Also, community models have their challenges. For example, how is the model verified once modifications have been made? Is there a series of verification examples? Is there a single version or have many “branches” been developed over the years? If there are many branches or versions, how are they managed? Who is responsible for maintaining the code and designing the development and sustainability plan? What are the main issues faced by a user (new or experienced)?

Authors Response

We have included further descriptions to the software development and sustainability plan for ParFlow such as new formulations of both kinematic and diffusive wave approximations, and advanced parallelization support (GPU’S and heterogeneous compute architectures). See Lines 967-969 in the revised manuscript.

Sources to all versions and/or releases of the code has been provided in the “code availability and data policy” section where code developers and contributors can be found. This is not included in the main text to prevent redundancy. See Lines 984-986 in the revised manuscript.

ParFlow’s maintenance and development and sustainability plans are designed by group of code developers and scientists from various institutions listed in Lines 131-135 in the revised manuscript.

Lastly, ParFlow has a clear, rigorous verification procedure to make sure that any changes checked in do not “break” previous developments. This ensures numerical accuracy and backwards compatibility. The full suite of test cases is automatically re-run before any submitted change can be considered for merging with the master branch of the code. The number of branches/forks cannot be controlled in any open source (or community) code, but any contributions to the master branch are exhaustively vetted before being pushed out to users. A note explaining this procedure has been added (Lines 958-969).

Tables and Figures

Table 2 provides an overview of coupling studies but with very little information and one has to refer to the individual publications to have a better understanding of these simulations (and ParFlow’s capabilities). In that table,

- The simulation scale is not clear since there are mentions of watershed and catchment but it is not clear what size they are. There is also a mention of regional scale but no indication on how it is different from catchment or watershed. I suggest that some information on the size of the model (for example the area and perhaps the depth) be given.

Authors Response

Table 2 has been revised to define the sizes (i.e. lateral and vertical dimensions) of catchment, watershed, regional scale as used in the original articles mentioned in the manuscript (Lines 1125-1127).

- It would be informative, for a potential user, to indicate which studies are conceptual (e.g. model development, numerical methods) and which are application to real systems, with a mention if there was a model calibration to observations.

Authors Response

Table 2 has been revised to indicate whether the original study was model development and if there was a model calibration to observations.
Figure 7 is not referenced in the text.

Authors Response
The paper has been revised to reference Figure 7 in the text. See Line 902 in the revised manuscript.

Symbols and equations
The symbols used in the equations have to be checked for consistency. There are several instances where the same letter or symbol designates different quantities and cases where the same quantity is identified with a different symbol (one example is hydraulic conductivity). Also, some variables (one example is porosity) are defined more than once. I am not providing an exhaustive list but some examples are:
• Equation 2: x is not defined
Authors Response
Equation 2 has been revised to define all variables appropriately (Line 167).

• In equation 2, p is pressure head but it is hydraulic head in equation 4
Authors Response
The symbols for pressure head and hydraulic head in Equations 2 and 4 have been revised. Different symbols have been used (Lines 167 and 186).

• Units for q_s in equation 1 are given as L^3T^{-1}, which is not consistent with the units for equation 1.
• q_s is used in both equations 1 and 5 but it is not the same quantity since the units are different in the two equations.
Authors Response
The units of q_s in Equations 1 and 5 have been revised to be equal (Line 172 and 207).

• Equation 5 could be deleted and replaced by equation 9
Authors Response
Equation 5 is a lead to equation 9 so we believe it does not make it less important including it.

• Equations presented in section 5 should be carefully reviewed because they have obviously been copied from other documents and have not been checked for consistency with respect to the ParFlow equations presented in section 2.
Authors Response
Equations in section 5 have been revised for consistency in the equations where appropriate.

Anonymous Referee #2 Comment
Writing
Careful proofreading is required because there are several instances where words are missing or where a sentence or expression is not clear. I am not providing an exhaustive list but some illustrative examples in the beginning of the paper are:
• Line 57: “vadose flow”. Should be something like vadose zone flow.
Authors Response
Line 57 is revised to include “zone” to vadose flow. See Line 57 in the revised manuscript.

• Lines 58-59: “process domains”. Not sure what process domains are.
Authors Response
Lines 58-59 have been revised. See Line 59 in the revised manuscript.
• Lines 62-63: “hydraulically-linked interconnected” is redundant
Authors Response
Text in Line 63 are revised to eliminate redundancy. The phrase “hydraulically-connected” is used.

• Line 64: “feedback between the components”. Components is not defined and it is not clear what it refers to.
Authors Response
Components represented surface and subsurface flow systems, and the text has been revised as such in Line 64.

• Lines 75-76: “surface, unsaturated, and groundwater flow”. There should not be any distinction between unsaturated flow and groundwater flow. Flow in the unsaturated zone is groundwater flow.
Authors Response
“Unsaturated” has been removed. See Line 76 in the revised manuscript.

• Line 77: “surface and overland flow”. Is surface flow different from overland flow? The paper uses both terms without specifying if they are synonyms or represent different flow processes (which this sentence is suggesting). The paper should be checked for consistency in using surface and/or overland flow.
Authors Response
The paper is revised to use surface or overland flow as synonyms.
Simulating Coupled Surface-Subsurface Flows with ParFlow v3.5.0: Capabilities, applications, and ongoing development of an open-source, massively parallel, integrated hydrologic model

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Surface and subsurface flow constitute a naturally linked hydrologic continuum that has not traditionally been simulated in an integrated fashion. Recognizing the interactions between these systems has encouraged the development of integrated hydrologic models (IHMs) capable of treating surface and subsurface systems as a single integrated resource. IHMs is dynamically evolving with improvement in technology and the extent of their current capabilities are often only known to the developers and not general users. This article provides an overview of the core functionality, capability, applications, and ongoing development of one open-source IHM, ParFlow. ParFlow is a parallel, integrated, hydrologic model that simulates surface and subsurface flows. ParFlow solves Richards’ equation for three-dimensional variably saturated groundwater flow and the two-dimensional kinematic wave approximation of the shallow water equations for overland flow. The model employs a conservative centered finite difference scheme and a conservative finite volume method for subsurface flow and transport, respectively. ParFlow uses multigrid preconditioned Krylov and Newton-Krylov methods to solve the linear and nonlinear systems within each time step of the flow simulations. The code has demonstrated very efficient parallel solution capabilities. ParFlow has been coupled to geochemical reaction, land surface (e.g. Common Land Model), and atmospheric models to study the interactions among the subsurface, land surface, and the atmosphere systems across different spatial scales. This overview focuses on the current capabilities of the code, the core simulation engine, and the primary couplings of the subsurface model to other codes, taking a high-level perspective.
1. Introduction

Surface and subsurface (unsaturated and saturated zones) water are connected components of a hydrologic continuum (Kumar et al., 2009). The recognition that flow systems (i.e. surface and subsurface) are a single integrated resource has stimulated the development of integrated hydrologic models (IHMs), which include codes like ParFlow (Ashby and Falgout, 1996; Kollet and Maxwell, 2006), HydroGeoSphere (Therrien and Sudicky, 1996), PIHM (Kumar, 2009), and CATHY (Camporese et al., 2010). These codes explicitly simulate different hydrological processes such as feedbacks between processes that affect the timing and rates of evapotranspiration, vadose zone flow, surface runoff and groundwater interactions. That is, IHMs are designed specifically to include the interactions between traditionally incompatible process flow domains (e.g. groundwater and land surface flow) (Engdahl and Maxwell, 2015). Most IHMs adopt a similar, physically-based approach to describe watershed dynamics where the governing equations of three–dimensional variably saturated subsurface flow are coupled to shallow water equations for surface runoff. The advantage of the coupled approach is that it allows hydraulically-linked interconnected groundwater–surface water systems to evolve dynamically, and for natural feedbacks between the components-systems to develop (Sulis et al., 2010; Maxwell et al., 2011; Weill et al., 2011; Williams and Maxwell, 2011; Simmer et al., 2015). A large body of literature now exists presenting applications of the various IHMs to solve hydrologic questions. Each model has its own technical documentation, but the individual development, maintenance, and sustainability efforts differ between tools. Some IHMs represent commercial investments and...
others are community, open-sourced projects, but all are dynamically evolving as technology improves and new features are added. Consequently, it can be difficult to answer the question of “what exactly can this IHM do today” without navigating dense user documentation. The purpose of this manuscript is to provide a current review of the functions, capabilities, and ongoing development of one of the open-source integrated models, ParFlow, in a format that is more accessible to a broad audience than a user manual or articles detailing specific applications of the model.

ParFlow is a parallel integrated hydrologic model that simulates surface, unsaturated, and groundwater flow (Maxwell et al., 2016). ParFlow computes fluxes through the subsurface, as well as interactions with above-ground or surface (overland) flow: all driven by gradients in hydraulic head. Richards’ equation is employed to simulate variably saturated three-dimensional groundwater flow (Richards, 1931). Overland flow can be generated by saturation or infiltration excess using a free surface overland flow boundary condition combined with Manning’s equation and the kinematic wave formulations of the dynamic wave equation (Kollet and Maxwell, 2006). ParFlow solves these governing equations employing either a fully coupled or integrated approach where surface and subsurface flows are solved simultaneously the Richards’ equation in three-dimensional form (Gilbert and Maxwell, 2016), or an indirect approach where the different components can be partitioned and flows in only one of the systems (surface or subsurface flows) is solved. The integrated approach allows for dynamic evolution of the interconnectivity between the surface water and groundwater systems. This interconnection depends only on the properties of the physical system and governing equations. An indirect approach permits partitioning of the...
flow components i.e. water and mass fluxes between surface and subsurface systems. The flow components can be solved sequentially. For the groundwater flow solution, ParFlow makes use of an implicit backward Euler scheme in time, and a cell-centered finite-difference scheme in space (Woodward, 1998). An upwind finite-volume scheme in space and an implicit backward Euler scheme in time is used for the overland flow component (Maxwell et al., 2007). ParFlow uses Krylov linear solvers with multigrid preconditioners for the flow equations along with a Newton method for the nonlinearities in the variably saturated flow system (Ashby and Falgout, 1996; Jones and Woodward, 2001). ParFlow’s physically based approach requires a number of parameterizations e.g. subsurface hydraulic properties, such as porosity, the saturated hydraulic conductivity, and the pressure-saturation relationship parameters (relative permeability), etc. (Kollet and Maxwell, 2008a; Kollet et al., 2010).

ParFlow is well documented and has been applied to surface and subsurface flow problems including simulating the dynamic nature of groundwater and surface-subsurface interconnectivity in large domains (e.g. over 600 km²) (e.g. Kollet and Maxwell, 2008; Ferguson and Maxwell, 2012; Condon et al., 2013; Condon and Maxwell, 2014), small catchments (e.g. approximately 30 km²) (Ashby et al., 1994; Kollet and Maxwell, 2006; Engdahl et al., 2016), complex terrain with highly heterogenous subsurface permeability such as the Rocky Mountain National Park, Colorado, United States (e.g. Engdahl and Maxwell, 2015; Kollet et al., 2017), large watersheds (Abu-El-Sha’r and Rihani, 2007; Kollet et al., 2010), continental scale flows (Condon et al., 2015; Maxwell et al., 2015) and even subsurface–surface and –atmospheric coupling (Maxwell et al., 2011; Williams and Maxwell, 2011; Williams et al., 2013; Gasper et al., 2014; Shrestha et al.,
Evidences from these studies suggest ParFlow produce accurate results in simulating flows in surface-subsurface systems in watersheds i.e. the code possesses the capability of performing simulations that accurately represent the behaviors of natural systems on which models are based.

The rest of the paper is organized as follows: We provide a brief history of ParFlow’s development in Section 1.1. In Section 2, we describe the core functionality of the code, i.e. the primary functions and the model equations and grid type used by ParFlow. Section 3 covers equation discretization and solvers (e.g. inexact Newton-Krylov, the ParFlow Multigrid (PFMG) preconditioner, and the Multigrid-Preconditioned Conjugate Gradient (MGCG) method) used in ParFlow. Examples of parallel scaling and performance efficiency of ParFlow are revisited in Section 4. The coupling capabilities of ParFlow, with other atmospheric, land surface, and subsurface models are shown in Section 5. We provide a summary and discussion, future directions to the development of ParFlow, and give some concluding remarks in Section 6.

1.1 Development History

ParFlow development commenced as part of an effort to develop an open-source, object-oriented, parallel watershed flow model initiated by scientists from the Center for Applied Scientific Computing (CASC), Environmental Programs, and the Environmental Protection Department at the Lawrence Livermore National Laboratory (LLNL) in the mid–1990s. ParFlow was born out of this effort to address the need for a code that combines fast, nonlinear solution schemes with massively parallel processing power, and its development continues today (e.g. Ashby et al., 1993; Smith et al., 1995; Woodward, 1998; Maxwell and Miller, 2005; Kollet and
Maxwell, 2008; Rihani et al., 2010; Simmer et al., 2015). ParFlow, is now a collaborative effort between numerous institutions including Colorado School of Mines, Research Center Jülich, University of Bonn, Washington State University, the University of Arizona, and Lawrence Livermore National Laboratory, and its working base and development community continues to expand.

ParFlow was originally developed for modeling saturated fluid flow and chemical transport in three-dimensional heterogeneous media. Over the past few decades, ParFlow underwent several modifications and expansions (i.e. additional features and capabilities have been implemented) and has seen an exponential growth of applications. For example, a two-dimensional distributed overland flow simulator (surface water component) was implemented into ParFlow (Kollet and Maxwell, 2006) to simulate interaction between surface and subsurface flows. Such additional implementations have resulted in improved numerical methods in the code. In The code’s applicability continues to evolve, for example, in recent times, ParFlow has been used in several coupling studies, with subsurface, land surface, and atmospheric models to include physical processes at the land surface (Maxwell and Miller, 2005; Maxwell et al., 2007, 2011; Kollet, 2009; Williams and Maxwell, 2011; Valcke et al., 2012; Valcke, 2013; Shrestha et al., 2014; Beisman et al., 2015) across different spatial scales and resolutions (Kollet and Maxwell, 2008; Condon and Maxwell, 2015; Maxwell et al., 2015). Also, a terrain following mesh formulation has been implemented (Maxwell, 2013) that allows ParFlow to handle problems with fine space discretization near the ground surface that comes with variable vertical discretization flexibility.
which offer modelers the advantage to increase the resolution of the shallow soil layers (these are discussed in detail below).

2. Core Functionality of ParFlow

The core functionality of the ParFlow model is the solution of three-dimensional variably saturated groundwater flow in heterogeneous porous media ranging from simple domains with minimal topography and/or heterogeneity to highly resolved continental-scale catchments (Jones and Woodward, 2001; Maxwell and Miller, 2005; Kollet and Maxwell, 2008; Maxwell, 2013). Within this range of complexity, the ParFlow model can operate in three different modes: 1). variably saturated; 2). steady-state saturated; and 3). integrated–watershed flows; however, all these modes share a common sparse coefficient matrix solution framework.

2.1 Variably Saturated Flow

ParFlow can operate in variably saturated mode using the well-known, mixed form of Richards’ equation (Celia et al., 1990). The mixed form of Richards’ equation implemented in ParFlow is:

\[ S_s S_w(p) \frac{\partial p}{\partial t} + \phi \frac{\partial (S_w(p))}{\partial t} = \nabla \cdot q + q_s, \quad (1) \]

\[ q = -k_s(\mathbf{x})k_r(p)\nabla(p - z), \quad (2) \]

where \( S_s \) is the specific storage coefficient \([\text{L}^{-1}]\), \( S_w \) is the relative saturation \([-\] as a function of pressure head \( p \) of the fluid/water \([\text{L}]\), \( t \) is time \([\text{T}]\), \( \phi \) is the porosity of the medium \([-\]), \( q \) is
the specific volumetric (Darcy) flux \([LT^{-1}]\), \(k_s(x)\) is the saturated hydraulic conductivity tensor \([LT^{-1}]\), \(k_r\) is the relative permeability \([-\)] which is a function of pressure head, \(q_s\) is the general source/sink term \([L^2T^{-1}]\) (includes wells and surface fluxes e.g. evaporation and transpiration), and \(z\) is depth below the surface \([L]\). The Richards’ equation assumes that the air phase is infinitely mobile (Richards, 1931). ParFlow has been used to numerically simulate river-aquifer exchange (free-surface flow and subsurface flow), (Frei et al., 2009), and highly heterogenous problems under variably-saturated flow conditions (Woodward, 1998; Jones and Woodward, 2001; Kollet et al., 2010). Under saturated conditions e.g. simulating linear groundwater movement under assumed predevelopment conditions, the steady-state saturated mode can be used.

2.2 Steady-State Saturated Flow

The most basic operational mode is the solution of the steady state, fully saturated groundwater flow equation:

\[ \nabla \cdot q - q_s = 0, \]

where \(q_s\) represents a general source/sink term e.g. wells \([L^2T^{-1}]\), \(q\) is the Darcy’ flux \([L^2T^{-1}]\) which is usually written as:

\[ q = -k_s \nabla \cdot P \]

where \(k_s\) is the saturated hydraulic conductivity \([LT^{-1}]\) and \(P\) represents the 3-D hydraulic head-potential \([L]\). ParFlow does include a direct solution option for the steady state saturated flow that is distinct from the transient solver. For example, ParFlow uses the solver “impes” under single-phase, fully saturated steady state condition relative to the variably saturated, transient mode where
Richards’ equation solver is used (Maxwell et al., 2016). When studying sophisticated or complex phenomena e.g. simulating fully coupled system (i.e. surface and subsurface flow), an overland flow boundary condition is employed.

2.3 Overland Flow

Surface water systems are connected to the subsurface, and these interactions are particularly important for rivers. However, these connections have been historically difficult to represent explicitly in numerical simulations. A common approach has been to use river routing codes, like HEC, and MODFLOW and its River Package to determine head in the river, which is then used as a boundary condition for the subsurface model. This approach prevents feedbacks between the two models, and a better representation of the physical processes in these kinds of problems is one of the motivations for IHMs. Overland flow is implemented in ParFlow as a two-dimensional kinematic wave equation approximation of the shallow water equations. The continuity equation for two-dimensional shallow overland flow is given as;

\[ \frac{\partial \psi_s}{\partial t} = \nabla \cdot (\bar{v} \psi_s) + q_s, \]  

(5)

where \( \bar{v} \) is the depth averaged velocity vector \([LT^{-1}]\), \( \psi_s \) is the surface ponding depth \([L]\), \( t \) is time \([T]\), and \( q_s \) is a general source/sink (e.g. precipitation rate) \([LT^{-1}]\). Ignoring the dynamic and diffusion terms results in the momentum equation

\[ S_{f,i} = S_{o,i}, \]

(6)
which is known as the kinematic wave approximation. The $S_{f,i}$ and $S_{o,i}$ represent the friction $[-]$ and bed slopes (gravity forcing term) $[-]$ slopes, respectively, where $i$ indicates $x -$ and $y -$ directions (also shown in equations 7 and 8) (Maxwell et al., 2015). Manning’s equation is used to generate a flow depth–discharge relationship:

\[ u_x = \frac{\sqrt{S_{f,x}}}{n} \psi_s^{2/3}, \text{ and} \]
\[ u_y = \frac{\sqrt{S_{f,y}}}{n} \psi_s^{2/3} \]  

where $n$ is the Manning’s roughness coefficient $[\text{TL}^{-1/3}]$. Flow of water out of overland flow simulation domain only occurs horizontally at an outlet which is controlled by specifying a type of boundary condition at the edge of the simulation domain. In a natural system, the outlet is usually taken as the region where a river enters another water body such as stream or a lake. ParFlow determines overland flow direction through the D4 flow routing approach. In a simulation domain, the D4 flow routing approach allows for flow to be assigned from a focal cell to only one neighboring cell accessed via the steepest or most vertical slope. The shallow overland flow formulation (equation 9) assumes that the flow depth is averaged-vertically and neglects a vertical change in momentum in the column of surface water. To account for vertical flow (from the surface to the subsurface or subsurface to the surface into the subsurface from the surface), a formulation that couples the system of equations through a boundary condition at the land surface becomes necessary. Equation (5) can be modified to include an exchange rate with the subsurface, $q_e$, as:

\[ \frac{\partial \psi_s}{\partial t} = \nabla \cdot (\hat{\nu} \psi_s) + q_s + q_e \]  

(9)
which is common in other IHMs. In ParFlow, the overland flow equations are coupled directly to Richards’ equation at the top boundary cell under saturated conditions. Conditions of continuity of pressure (i.e. the pressures of the subsurface and surface domains are equal right at the ground surface) and flux at the top cell of the boundary between the subsurface and surface systems are assigned. Figure 1 is provided demonstrating continuity of pressure at the ground surface for flow from the surface into the subsurface. This assignment is done by setting pressure–head, in equation (1) equal to the vertically–averaged surface pressure, $\psi_s$;

$$p = \psi_s = \psi,$$  

(10)

and the flux, $q_e$ equal to the specified boundary conditions (e.g. Neumann or Dirichlet type). For example, if Neumann type boundary conditions are specified, which are given as;

$$q_{BC} = -k_s k_r \nabla (\psi - z)$$  

(11)

and one solves for the flux term in equation (10), the result is;

$$q_e = \frac{\partial \|\psi, 0\|}{\partial t} - \nabla \|\psi, 0\| - q_s$$  

(12)

where the $\|\psi, 0\|$ operator is defined as the greater of the quantities, $\psi$ and $0$. Substituting equation (12) for the boundary condition in equation (11), requiring the aforementioned flux continuity $q_{BC} = q_e$, leads to

$$-k_s k_r \nabla (\psi - z) = \frac{\partial \|\psi, 0\|}{\partial t} - \nabla (\|\psi, 0\|) - q_s$$  

(13)

Equation (13) shows that the surface water equations are represented as a boundary condition to the Richards’ equation. That is, the boundary condition links flow processes in the subsurface with those at the land surface. This boundary condition eliminates the exchange flux and accounts for
the movement of the free surface of ponded water at the land surface (Kollet and Maxwell, 2006;
Williams and Maxwell, 2011).

Many IHMs couple subsurface and surface flows making use of the exchange flux, $q_e$
model. The exchange flux between the domains (the surface and the subsurface) depends on
hydraulic conductivity and the gradient across some interface where indirect coupling is used
(VanderKwaak, 1999; Panday and Huyakorn, 2004). The exchange flux concept gives a general
formulation of a single set of coupled surface-subsurface equations. The exchange flux term, $q_e$
may be included in the shallow overland flow continuity equation as the exchange rate term with
the subsurface (equation 9) in a coupled system (Kollet and Maxwell, 2006).

Figure. 1 Caption: Coupled surface and subsurface flow systems. Note in this figure the physical
system is represented on the left and a schematic of the overland flow boundary condition
(continuity of pressure and flux at the ground surface) is on the right. The equation, $p = \psi_s = \psi$
in Figure. 1 signifies that the vertically averaged surface pressure and subsurface pressure head are
equal right at the land surface.

2.4 Multi-Phase Flow and Transport Equations

Most applications of the code have reflected ParFlow’s core functionality as a single-phase
flow solver, but there are also embedded capabilities for multi-phase flow of immiscible fluids and
solute transport. Multi–phase systems are distinguished from single–phase systems by the presence
of one or more interfaces separating the phases, with moving boundaries between phases. The flow
equations that are solved in multi-phase systems in a porous medium comprise a set of mass balance and momentum equations. The equations are given by:

\[
\frac{\partial}{\partial t} (\phi \rho_i S_i) + \nabla \cdot (\phi \rho_i S_i \vec{v}_i) - \rho_i Q_i = 0, \quad (14)
\]

\[
\phi S_i \vec{v}_i + \lambda_i \cdot (\nabla p_i - \rho_i \vec{g}) = 0, \quad (15)
\]

where \(i = 1, \ldots, n\) denotes a given phase (such as air or water). In these equations, \(\phi\) is the intrinsic medium porosity \([-]\) which explains the fluid capacity of the porous medium, and for each phase, \(i, S_i(\vec{x}, t)\) is the relative saturation \([-]\) which indicates the content of phase \(i\) in the porous medium, \(\vec{v}_i(\vec{x}, t)\) represent Darcy velocity vector \([LT^{-1}]\), \(Q_i(\vec{x}, t)\) stands for source/sink term \([T^{-1}]\), \(p_i(\vec{x}, t)\) is the average pressure \([ML^{-1}T^{-2}]\), \(\rho_i(\vec{x}, t)\) is the mass density \([ML^{-3}]\), \(\lambda_i\) is the mobility \([L^3TM^{-1}]\), \(\vec{g}\) is the gravity vector \([LT^{-2}]\), and \(\vec{x}\) and \(t\) represent space vectors and the time respectively. ParFlow solves for the pressures on a discrete mesh and uses a time-stepping algorithm based on a mass conservative backward Euler scheme and spatial discretization (a finite volume method). ParFlow’s multi-phase flow capability has not been applied in major studies, however, this capability is also available for testing (Ashby et al., 1993; Tompson et al., 1994; Falgout et al., 1999; Maxwell et al., 2016).

The transport equations included in the ParFlow package describe mass conservation in a convective flow (no diffusion) with degradation effects and adsorption included along with extraction and injection wells (Beisman et al., 2015; Maxwell et al., 2016). The transport equation is defined as follows:
\[
\left( \frac{\partial}{\partial t} (\phi c_{i,j}) + \lambda_j \phi c_{i,j} \right) + \nabla \cdot (c_{i,j}\vec{v}) = -\left( \frac{\partial}{\partial t} (1 - \phi) \rho_s F_{i,j} \right) + \lambda_i (1 - \phi) \rho_s F_{i,j} + \nabla \cdot \left( c_{i,j} \vec{v} \right) = -\left( \frac{\partial}{\partial t} ((1 - \phi) \rho_s F_{i,j}) + \lambda_i (1 - \phi) \rho_s F_{i,j} \right) + \lambda_j \phi c_{i,j}
\]

where \( \phi \) denotes the porosity of the flow medium \([-]\), \( c_{i,j}(\vec{x}, t) \) represents concentration fraction of contaminant \([-]\), \( \vec{v}(\vec{x}, t) \) is the Darcy velocity vector \([LT^{-1}]\), \( \lambda_i \) is degradation rate \([T^{-1}]\), \( F_{i,j}(\vec{x}, t) \) is the mass concentration \([L^3M^{-1}]\), \( \rho_s(\vec{x}) \) is the density of the solid mass \([ML^{-3}]\), \( n_i \) is injection wells \([-]\), \( \gamma_{k}^{I;i}(t) \) is injection rate \([T^{-1}]\), \( \Omega_{k}^{I}(\vec{x}) \) represent the area of the injection well \([-]\), \( c_{i,j}^{-k}(\vec{x}, t) \) is the injected concentration fraction \([-]\), \( n_E \) is the extraction wells \([-]\), \( \gamma_{k}^{E;i}(t) \) is extraction rate \([T^{-1}]\), \( \Omega_{k}^{E}(\vec{x}) \) is an extraction well area \([-]\), \( i = 0, ..., n_p-1 \) \( (n_p \in \{1, 2, 3\}) \) is the number of phases, \( j = 0, ..., n_c - 1 \) represents the number of contaminants, \( c_{i,j} \) is the concentration of contaminant \( j \) in phase \( i \), \( k \) is hydraulic conductivity \([LT^{-1}]\), \( \chi_{k}^{I} \) is the characteristic function of an injection well region, and \( \chi_{k}^{E} \) is the characteristic function of an extraction well region. The mass concentration term, \( F_{i,j} \) is taken to be instantaneous in time and a linear function of contaminant concentration:

\[
F_{i,j} = K_{d;i} c_{i,j}
\]

where \( K_{d;i} \) is the distribution coefficient of the component \([L^3M^{-1}]\). The transport/advection equation or convective flow calculation performed by ParFlow offers a choice of a first-order explicit upwind scheme or a second-order explicit Godunov scheme. The advection calculations are discretized as boundary value problems for each primary dimension over each compute cell. The discretization is a fully-explicit, forward Euler first-order accurate in time approach. The implementation of a second-order explicit Godunov scheme (second-order advection scheme)
minimizes numerical dispersion and presents accurate computational process at these time scales than either an implicit or lower-order explicit scheme. Stability issue here is that the simulation timestep is restricted via the courant-Friedrichs-Lewy (CFL) condition, which demands that time steps are chosen small enough to ensure that mass not be transported more than one grid cell in a single timestep in order to maintain stability (Beisman, 2007).

2.5 Computational Grids

An accurate numerical approximation of a set of partial differential equations is strongly dependent on the simulation grid. Integrated hydrologic models can use unstructured or structured meshes for the discretization of the governing equations. The choice of grid type to adopt is problem-specific and often a subjective choice since the same domain can be represented in many ways, but there are some clear tradeoffs. For example, structured grid models, such as ParFlow, may be preferred to unstructured grid models because structured grids provide significant advantages in computational simplicity and speed, and are amenable to efficient parallelization (Durbin, 2002; Kumar et al., 2009; Osei-Kuffuor et al., 2014). ParFlow adopts a regular, structured grid specifically for its parallel performance. There are currently two regular grid formulations included in ParFlow, an orthogonal grid and a terrain-following formulation (TFG); both allow for variable vertical discretization (thickness over an entire layer) over the domain.

2.5.1 Orthogonal Grid

Orthogonal grids have many advantages, and many approaches are available to transform an irregular grid into an orthogonal grid such as conformal mapping. This mapping defines a
transformed set of partial differential equations using an elliptical system with “control functions”
determined in such a way that the generated grid would be either orthogonal or nearly orthogonal.
However, conformal mapping may not allow flexibility in the control of the grid node distribution,
which diminishes its usefulness with complex geometries (Mobley and Stewart, 1980; Haussling
and Coleman, 1981; Visbal and Knight, 1982; Ryskin and Leal, 1983; Allievi and Calisal, 1992;
Eca, 1996).

A Cartesian, regular, orthogonal grid formulation is implemented by default in ParFlow,
though some adaptive meshing capabilities are still included in the source code. For example,
layers within a simulation domain can be made to have varying thickness. The upper portion of
Figure 2 shows the standard way topography or any other non-rectangular domain boundaries are
represented in ParFlow. The domain limits, and any other internal boundaries, can be defined using
grid-independent triangulated irregular network (TIN) files that define a geometry, or a gridded
indicator file can be used to define geometric elements. ParFlow uses octree space partitioning
algorithm (a grid-based algorithm or mesh generators filled with structured grids) (Maxwell, 2013)
to depict complex structures/land surface representations (e.g. topography, watershed boundaries,
and different hydrologic facies) in three-dimensional space (Kollet et al., 2010). These land surface
features are mapped onto the orthogonal grid, and looping structures that encompass these irregular
shapes are constructed (Ashby et al., 1997). The grid cells above ground surface are inactive
(shown in upper region of Figure 2) and are stored in the solution vector but not included in the
solution.

2.5.2 Terrain Following Grid
The inactive portion of a watershed defined with an orthogonal grid can be quite large in complex watersheds with high-relief. In these cases, it is advantageous to use a grid that allows these regions to be omitted. ParFlow’s structured grid conforms to the topography via transformation by the terrain following grid formulation. This transform alters the form of Darcy’s law to incorporate a topographic slope component. For example, subsurface fluxes are computed separately in both $x$ and $y$ directions making use of the terrain following grid transform as:

$$q_x = K \sin(\theta_x) + K \frac{\partial p}{\partial y} \cos(\theta_x), \text{ and}$$

$$q_y = K \sin(\theta_y) + K \frac{\partial p}{\partial y} \cos(\theta_y)$$  \hspace{1cm} (18)

where $q_x$ and $q_y$ represent source/sink terms, such as fluxes, that include potential recharge flux at the ground surface [LT$^{-1}$], $p$ is the pressure head [L]; $K$ is the saturated hydraulic conductivity tensor, [LT$^{-1}$], $\theta$ is the local angle [\degree] of topographic slope, $S_x$ and $S_y$ in the $x$ and $y$ directions and may be presented as $\theta_x = \tan^{-1} S_x$ and $\theta_y = \tan^{-1} S_y$ respectively (Weill et al., 2009). The terrain following grid formulation comes handy when solving coupled surface and subsurface flows (Maxwell, 2013). The terrain following grid formulation uses the same surface slopes specified for overland flow to transform the grid, whereas the slopes specified in the orthogonal grid are only used for 2-D overland flow routing and do not impact the subsurface formulation (see Figure 2). Note that TIN files can still be used to deactivate portions of the transformed domain.
Figure 2 Caption: Representation of orthogonal (upper) and the terrain following (lower) grid formulations and schematics of the related finite difference dependences (left). The $i, j, k$ are the $x, y, z$ cell indices.

3. Equation Discretization and Solvers

The core of the ParFlow code is its library of numerical solvers. As noted above, in most cases, the temporal discretization of the governing equations uses an implicit (backward Euler) scheme; with cell-centered finite differences in spatial dimensions. Different components of this solution framework have been developed for the various operational modes of ParFlow including an inexact Newton-Krylov nonlinear solver (section 3.1), a multigrid algorithm (section 3.2), and a multigrid-preconditioned conjugate gradient (MGCG) solver in (section 3.3). The conditions, requirements, and constraints on the solvers depend on the specifics of the problem being solved, and some solvers tend to be more efficient (faster overall convergence) than others for a given problem. The core structure of these solvers and some of their implementation details are given below, with an emphasis on the main concepts behind each solver.

3.1 Newton–Krylov solver for Variably Saturated Flow

The cell-centered fully-implicit discretization scheme applied to Richards’ equation leads to a set of coupled discrete nonlinear equations that need to be solved at each time step, and, for variably saturated subsurface flow, ParFlow does this with the inexact Newton-Krylov method.
implemented in the KINSOL package (Hindmarsh et al., 2005; Collier et al., 2015). Newton-Krylov methods were initially utilized in the context of partial differential equations by (Brown and Saad, 1990). In the approach, coupled nonlinear system as a result of discretization of the partial differential equation is solved iteratively. Within each iteration, the nonlinear system is linearized via a Taylor expansion. After linearization, an iterative Krylov method is used to solve the resulting linear Jacobian system (Woodward, 1998; Osei-Kuffuor et al., 2014). For variably saturated subsurface flow, ParFlow uses the GMRES Krylov method (Saad and Schultz, 1986). Figure 3 is a flow chart of the solution technique ParFlow uses to provide approximate solutions to systems of nonlinear equations.

The benefit of this Newton-Krylov method is that the Krylov linear solver requires only matrix-vector products. Because the system matrix is the Jacobian of the nonlinear function, these matrix-vector products may be approximated by taking directional derivatives of the nonlinear function in the direction of the vector to be multiplied. This approximation is the main advantage of the Newton-Krylov approach as it removes the requirement for matrix entries in the linear solver. An inexact Newton method is derived from a Newton method by using an approximate linear solver at each nonlinear iteration, as is done in the Newton-Krylov method (Dembo et al., 1982; Dennis and Schabel, 1996). This approach takes advantage of the fact that when the nonlinear system is far from converged, the linear model used to update the solution is a poor
approximation. Thus, the convergence criteria of early linear system solve is relaxed. The tolerance required for solution of the linear system is decreased as the nonlinear function residuals approach zero. The convergence rate of the resulting nonlinear solver can be linear or quadratic, depending on the algorithm used. Through the KINSOL package, ParFlow can either use a constant tolerance factor or ones from (Eisenstat and Walker, 1996). Krylov methods can be very robust, but they can be slow to converge. As a result, it is often necessary to implement a preconditioner, or accelerator, for these solvers.

3.2 Multigrid Solver

Multigrid (MG) methods constitute a class of techniques or algorithms for solving differential equations (system of equations) using a hierarchy of discretization (Volker, 1987; Briggs et al., 2000). Multigrid algorithms are applied primarily to solve linear and nonlinear boundary value problems and can be used as either preconditioners or solvers. The most efficient method for preconditioning the linear systems in ParFlow is the ParFlow Multigrid algorithm (PFMG) (Ashby and Falgout, 1996; Jones and Woodward, 2001). Multigrid algorithms arise from discretization of elliptic partial differential equations (Briggs et al., 2000), and, in ideal cases, have convergence rates that do not depend on the problem size. In these cases, the number of iterations remains constant even as problems sizes grow large. Thus, the algorithm is algorithmically scalable. However, it may take longer to evaluate each iteration as problem sizes increase. As a result, ParFlow utilizes the highly efficient implementation of PFMG in the hypre library (Falgout and Yang, 2002).
For variably saturated subsurface flow, ParFlow uses the Newton-Krylov method coupled with a multigrid preconditioner to accurately solve for the water pressure (hydraulic head) in the subsurface and diagnoses the saturation field (which is used in determining the water table). (Woodward, 1998; Jones and Woodward, 2000, 2001; Kollet et al., 2010). The water table is calculated for computational cells having hydraulic heads above the bottom of the cells. Generally, a cell is saturated if the hydraulic head in the cell is above the node elevation (cell center) or the cell is unsaturated if the hydraulic head in the cell is below the node elevation. For saturated flow, ParFlow uses the conjugate gradient method also coupled with a multigrid method. It is important to note that subsurface flow systems are usually much larger radially than they are thick, so it is common for the computational grids to have highly anisotropic cell aspect ratios to balance the lateral and vertical discretization. Combined with anisotropy in the permeability field, these high aspect ratios produce numerical anisotropy in the problem, which can cause the multigrid algorithms to converge slowly (Jones and Woodward, 2001). To correct this problem, a semicoarsening strategy or algorithm is employed, where the grid is coarsened in one direction at a time. The direction chosen is the one with the smallest grid spacing i.e. the tightest coupling. In an instance where more than one direction has the same minimum spacing, then the algorithm chooses the direction in the order of $x$, followed by $y$, and then in $z$. To decide on how and when to terminate the coarsening algorithm, Ashby and Falgout (1996) determined that a semicoarsening down to a $(1 \times 1 \times 1)$ grid is ideal for groundwater problems.

3.3 Multigrid-Preconditioned Conjugate Gradient (MGCG)
ParFlow uses the multigrid-preconditioned conjugate gradient (CG) solver to solve the groundwater equations under steady-state, and fully saturated flow conditions (Ashby and Falgout, 1996). These problems are symmetric and positive definite, two properties for which the CG method was designed to target. While CG lends itself to efficient implementations, the number of iterations required to solve a system such as results from discretization of the saturated flow equation increases as the problem size grows. The PFMG multigrid algorithm is used as a preconditioner to combat this growth and results in an algorithm for which the number of iterations required to solve the system grows only minimally. See Ashby and Falgout (1996) for a detailed description of these solvers and the parallel implementation of the multigrid preconditioned CG method in ParFlow (Gasper et al., 2014; Osei-Kuffuor et al., 2014).

3.4 Preconditioned Newton-Krylov for Coupled Subsurface – Surface Flows

As discussed above, coupling between subsurface and surface or overland flow in ParFlow is activated by specifying an overland boundary condition at the top surface of the computational domain, but this mode of coupling allows for activation and deactivation of the overland boundary condition during simulations where ponding or drying occur. Thus, surface–subsurface coupling can occur anywhere in the domain during a simulation and it can change dynamically during the simulation. Overland flow may occur by the Dunne or Horton mechanism depending on local dynamics. Overland flow routing is enabled when the subsurface cells are fully saturated. As discussed above, in ParFlow the coupling between the subsurface and surface flows is handled implicitly. ParFlow solves this implicit system with the inexact Newton-Krylov method.
described above. However, in this case, the preconditioning matrix is adjusted to include terms from the surface coupling. In the standard saturated or variably saturated case, the multigrid method is given the linear system matrix, or a symmetric version, resulting from discretization of the subsurface model. Because ParFlow uses a structured mesh, these matrices have a defined structure making their evaluation and application of multigrid straightforward. Due to varying topographic height of the surface boundary, where the surface coupling is enforced, the surface effects add non-structured entries in the linear system matrices. These entries increase complexity of the matrix entry evaluations and reduce effectiveness of the multigrid preconditioner. In this case, the matrix-vector products are most effectively performed through computation of the linear system entries, rather than the finite difference approximation to the directional derivative. For the preconditioning, surface couplings are only included if they model flow between cells at the same vertical height i.e. in situations where overland flow boundary conditions are imposed or activated. This restriction maintains the structured property of the preconditioning matrix while still including much of the surface coupling in the preconditioner. Both these adjustments led to considerable speedup in coupled simulations (Osei-Kuffuor et al., 2014).

4. Parallel Performance Efficiency

Scaling efficiency metrics offer a quantitative method for evaluating the performance of any parallel model. Good scaling generally means that the efficiency of the code is maintained as the solution of the system of equations is distributed onto more processors or as the problem resolution is refined and processing resources are added. Scalability can depend on the problem
size, the processor number, the computing environment, and the inherent capabilities of the computational platform used e.g. choice of a solver. The performance of ParFlow (or any parallel code) is typically determined through weak and strong scaling (Gustafson, 1988). Weak scaling involves the measurement of code’s efficiency in solving problems of increasing size (i.e. describes how the solution time change with change in the number of processors for a fixed problem size per processor). In weak scaling, the simulation time should remain constant, as the size of the problem and number of processing elements grow such that the same amount of work is conducted on each processing element. Following Gustafson (1988), scaled parallel efficiency is given by:

\[ E(n, p) = \frac{T(n,1)}{T(pn,p)} \]  

(19)

where \( E(n, p) \) denotes parallel efficiency, \( T \) represents the run time as a function of the problem size \( n \), which is spread across several processors \( p \). Parallel code is said to be perfectly efficient if \( E(n, p) = 1 \), and the efficiency decreases as \( E(n, p) \) approaches 0. Generally, parallel efficiency decreases with increasing processor number as communication overhead between nodes/processors becomes the limiting factor.

Strong scaling describes the measurement of how much the simulation or solution time changes with the number of processors for a given problem of fixed total size (Amdahl, 1967). In strong scaling, a fixed size task is solved on a growing number of processors, and the associated time needed for the model to compute the solution is determined (Woodward, 1998; Jones and Woodward, 2000). If the computational time decreases linearly with the processor number, a
perfect parallel efficiency, \( E = 1 \) results. The value of \( E \) is determined using equation (19). ParFlow has been shown to have excellent parallel performance efficiency, even for large problem sizes and processor counts (see Table 1) (Ashby and Falgout, 1996; Kollet and Maxwell, 2006). In situations where ParFlow works in conjunction with or coupled to other subsurface, land surface or atmospheric models (see Section 5) i.e. increased computational complexity by adding different components or processes, improved computational time may not only depend on ParFlow. The computational cost of such an integrated model is extremely difficult to predict because of the nonlinear nature of the system. The solution time may depend on number of factors including the number of degrees of freedom, the heterogeneity of the parameters, which processes are active (e.g. snow accumulation compared to nonlinear snowmelt processes in land surface model or the switching on or off of the overland flow routing in ParFlow). The only way to know how fast a specific problem will run is to try that problem. Many of the studies presented in Table 1 include computational times for problems with different complexities where ParFlow was used. In a scaling study with ParFlow, Maxwell (2013)—examined the relative performance of preconditioning the coupled variably saturated subsurface and surface flow system with the symmetric portion or full matrix for the system. Both options use ParFlow’s multigrid preconditioner. Solver performance was demonstrated by combining the analytical Jacobian and the non-symmetric linear preconditioner. The study showed that the non-symmetric linear preconditioner presents faster computational times and efficient scaling gains. A section of the study results is reproduced in Table 1, in addition to other scaling studies demonstrating ParFlow’s parallel efficiency. This tradeoff was also examined in Jones and Woodward (2000).
It is worth noting that large and/or complex problem sizes (e.g., simulating a large heterogeneous domain size with over 8.1 billion unknowns) will always take time to solve directly, but the approach for setting up a problem depends on the specific problem being modeled. Even for one specific kind of model there may be multiple workflows and how to model such complexity becomes sole responsibility of the modeler. The studies involving ParFlow outlined in Table 1 provide a wealth of knowledge regarding domain setup for problems of different complexities. Since these are all specific applications, their information will likely be very useful to modelers trying to build a new domain during the setup and planning phases.

Table 1: Details for the various parallel scaling studies conducted using ParFlow.

5. Coupling

Different integrated models including atmospheric or weather prediction models (e.g., Weather Research Forecasting Model, Advanced Regional Prediction System, Consortium for Small-Scale Modeling), land surface models (e.g., Common Land Model, Noah Land Surface Model), and a subsurface model (e.g., CruchFlow) have been coupled with ParFlow to simulate a variety of coupled earth system effects (see Figure 4(a)). Coupling between ParFlow and other integrated models was performed to better understand the physical processes that occur at the interfaces between the deeper subsurface and ground surface, and between the ground surface and the atmosphere. None of the individual models can achieve this on their own because ParFlow cannot
account for land surface processes (e.g. evaporation), and atmospheric and land surface models
generally do not simulate deeper subsurface flows (Ren and Xue, 2004; Chow et al., 2006;
Beisman, 2007; Maxwell et al., 2007; Shi et al., 2014). Model coupling can be achieved either via
“offline coupling” where models involved in the coupling process are run sequentially and
interactions between them is one–way (i.e. information is only transmitted from one model to the
other) or “online” where they interact and feedback mechanisms among components are
represented (Meehl et al., 2005; Valcke et al., 2009). Each of the coupled models uses its own
solver for the physical system it is solving, then information is passed between the models. As
long as each model exhibits good parallel performance, this approach still allows for simulations
at very high resolution, with a large number of processes (Beven, 2004; Ferguson and Maxwell,
2010; Shen and Phanikumar, 2010; Shi et al., 2014). This section focuses on the major couplings
between ParFlow and other codes. We point out specific functions of the individual models as
stand–alone codes that are relevant to the coupling process. In addition, information about the role
or contribution of each model at the coupling interface (see Figure 4(b)) that connects with
ParFlow are presented (Figure 5 shows the communication network of the coupled models). We
discuss couplings between ParFlow and its land surface model (a modified version of the original
Common Land Model introduced by Dai et al., (2003)), Consortium for Small-Scale Modeling
(COSMO), Weather Research Forecasting Model, Advanced Regional Prediction System, and
CruchFlow in sections 5.1, 5.2, 5.3, 5.4, and 5.5 respectively.

Figure 4(a) Caption: A pictorial description of the relevant physical environmental features and
model coupling. CLM represents the Community Land Model, a stand-alone Land Surface Model
via which ParFlow couples’ COSMO. The modified version of CLM by Dai et al., (2003) and is not shown in Figure 4(a) because it is a module only for ParFlow, not really a stand-alone LSM any longer.

Figure 4(b) Caption: Schematic showing information transmission at the coupling interface. PF, LSM, and ATM indicate the portions of the physical system simulated by ParFlow, Land Surface Models, and Atmospheric Models respectively. The downward and upward arrows indicate the directions of information transmission between adjacent models. Note: Coupling between ParFlow and CrunchFlow (not shown) occur within the subsurface.

5.1 ParFlow–Common Land Model (PF.CLM)

The Common Land Model (CLM) is a land surface model designed to complete land-water-energy balance at the land surface (Dai et al., 2003). CLM parameterizes the moisture, energy and momentum balances at the land surface and includes a variety of customizable land surface characteristics and modules, including land surface type (land cover type, soil texture, and soil color), vegetation and soil properties (e.g. canopy roughness, zero-plane displacement, leaf dimension, rooting depths, specific heat capacity of dry soil, thermal conductivity of dry soil, porosity), optical properties (e.g. albedos of thick canopy), and physiological properties related to the functioning of the photosynthesis-conductance model (e.g. green leaf area, dead leaf, and stem area indices). A combination of numerical schemes is employed to solve the governing equations. CLM uses a time integration scheme which proceeds by a split-hybrid approach, where the solution procedure is split into “energy balance” and “water balance” phases in a very modularized structure
The CLM described here and as incorporated in ParFlow is a modified version of the original CLM introduced by Dai et al., (2003) though the original version was coupled to ParFlow in previous model applications (e.g. Maxwell and Miller, 2005). The current coupled model, PF.CLM consist of ParFlow incorporated with land surface model Jefferson et al., (2015), (2017), and Jefferson and Maxwell, (2015). This results in the coupled model, PF.CLM i.e. ParFlow with its own land surface model. The modified CLM is composed of a series of land surface modules that are called as a subroutine within ParFlow to compute energy and water fluxes (e.g. evaporation and transpiration) to and out of the soil. For example, the modified CLM computes bare ground surface evaporative flux, $E_{gr}$ as

$$E_{gr} = -\beta \rho_a u_* q_*$$

(20)

where $\beta$ (dimensionless) denotes soil resistance factor, $\rho_a$ represents air density [ML$^{-3}$], $u_*$ represents friction velocity [LT$^{-1}$], and $q_*$ (dimensionless) stands for humidity scaling parameter (Jefferson and Maxwell, 2015). Evapotranspiration for vegetated land surface, $E_{veg}$ is computed as

$$E_{veg} = [R_{pp, dry} + L_w] L_{SAI} \left[ \frac{\rho_a}{r_b} (q_{sat} - q_{af}) \right]$$

(21)

where $r_b$ is the air density boundary resistance factor [LT$^{-1}$], $q_{sat}$ (dimensionless) is saturated humidity at the land surface, and $q_{af}$ (dimensionless) is the canopy humidity. Combination of $q_{sat}$ and $q_{af}$ forms the potential evapotranspiration. The potential evapotranspiration is divided into transpiration $R_{pp, dry}$ (dimensionless) which depends on the dry fraction of the canopy, and evaporation from foliage covered by water $L_w$ (dimensionless). $L_{SAI}$ (dimensionless) is summation.
of the leaf and stem area indices which estimates the total surface from which evaporation can occur. A detailed description of the equations CLM of PF.CLM uses can be found in Jefferson et al., (2015), (2017), and Jefferson and Maxwell, (2015).

The coupled model PF.CLM simulates variably saturated subsurface flow, surface or overland flow, and above-ground processes. PF.CLM was developed prior to the current community land model (see section 5.2), and the module structure of the current and early versions are different. PF.CLM has been updated over the years to improve its capabilities. PF.CLM was first done in the early 2000’s, as an undiversified, a column proof-of-concept model, where data or message was transmitted between the coupled models via input/output files (Maxwell and Miller, 2005). Later, PF.CLM was presented in a distributed or diversified approach with a parallel input/output file structure where CLM is called as a set sequence of steps within ParFlow (Kollet and Maxwell, 2008a). These modifications, for example, were done to incorporate subsurface pressure values from ParFlow into chosen computations (Jefferson and Maxwell, 2015). These, to some extent differentiate the modified version (PF.CLM) from the original CLM by Dai et al., (2003). Within the coupled PF.CLM, ParFlow solves the governing equations for overland and subsurface flow systems and the CLM modules add the energy balance and mass fluxes from the soil, canopy, and root zone that can occur (i.e. interception, evapotranspiration etc.) (Maxwell and Miller, 2005)(Jefferson and Maxwell, 2015).

At the coupling interface where the models overlap and undergo online communication (see Figure 4(b)), ParFlow calculates and passes soil moisture as well as pressure heads of the subsurface to CLM, and CLM calculates and transmits transpiration from
plants, canopy and ground surface evaporation, snow accumulation and melt, and infiltration from precipitation to ParFlow (Ferguson et al., 2016). In short, CLM does all canopy water balances and snow, but once the water through falls to the ground, or snow melts, ParFlow takes over and estimates the water balances via the nonlinear Richards’ equation. The coupled model, PF.CLM, has been shown to more accurately predict root-depth soil moisture compared to the uncoupled model i.e. stand-alone land surface model (CLM) with capability of computing near surface soil moisture. This increased accuracy results from the coupling of soil saturations determined by ParFlow and their impacts on other processes including runoff and infiltration (Kollet, 2009; Shrestha et al., 2014; Gebler et al., 2015; Gilbert and Maxwell, 2016). For example, (Maxwell and Miller, 2005) found that simulations of deeper soil saturation (more than 40cm) vary between PF.CLM and uncoupled models, with PF.CLM simulations closely matching the observed data. Table 2 contains summaries of studies conducted with ParFlow coupled to either the original version of CLM by (Dai et al., 2003) or modified CLM (ParFlow with land surface model).

5.1.1. ParFlowE–Common Land Model (ParFlowE[CLM])

It is well established that ParFlow in conjunction with CLM PF.CLM does perform well in estimating all canopy water and subsurface water balances (Maxwell and Miller, 2005; Mikkelson et al., 2013; Ferguson et al., 2016). ParFlow, as a component of the coupled model has been modified into a new parallel numerical model, ParFlowE to incorporate the more complete heat equation coupled to variably saturated flow. ParFlowE simulates coupling of terrestrial hydrologic and energy cycles i.e. coupled moisture, heat, and vapor transport in the subsurface. ParFlowE is based on the original version of ParFlow having identical solution schemes and coupling approach
with CLM. A coupled three-dimensional subsurface heat transport equation is implemented in ParFlowE using a cell-centered finite difference scheme in space and an implicit backward Euler differencing scheme in time. However, the solution algorithm employed in ParFlow is fully exploited in ParFlowE where the solution vector of the Newton-Krylov method was extended to two dimensions (Kollet et al., 2009). In some integrated and climate models, the convection term of subsurface heat flux and the effect of soil moisture on energy transport is neglected due to simplified parameterizations and computational limitations. However, both convection and conduction terms are considered in ParFlowE (Khorsandi et al., 2014). In ParFlowE, functional relationships (i.e. equations of state) are performed to relate density and viscosity to temperature and pressure, and thermal conductivity to saturation. That is, modeling thermal flows by relating these parameterizations in simulating heat flow is an essential component of ParFlowE. In coupling between ParFlowE and CLM, ParFlowE[CLM], the one-dimensional subsurface heat transport in the CLM is replaced by the three-dimensional heat transport equation including the process of convection of ParFlowE. CLM computes mass and energy balances at ground surface that lead to moisture fluxes and pass these fluxes to the subsurface moisture algorithm of ParFlowE[CLM]. These fluxes are used in computing subsurface moisture and temperature fields which are then passed back to the CLM.

5.2 ParFlow in the Terrestrial Systems Modeling Platform, TerrSysMP

ParFlow is part of the Terrestrial System Modeling Platform TerrSysMP, which comprise the nonhydrostatic fully compressible limited-area atmospheric prediction model, COSMO,
designed for both operational numerical weather prediction and various scientific applications on
the meso–β (horizontal scales of 20–200km) and meso–γ (horizontal scales of 2–20km) (Duniec
and Mazur, 2011; Levis and Jaeger, 2011; Bettems et al., 2015), and the Community Land Model
version 3.5 (CLM3.5). Currently, it is used in direct simulations of severe weather events triggered
by deep moist convection, including intense mesoscale convective complexes, prefrontal squall–
line storms, supercell thunderstorms, and heavy snowfall from wintertime mesocyclones. COSMO
solves nonhydrostatic, fully compressible hydro–thermodynamical equations in advection form
using the traditional finite difference method (Vogel et al., 2009; Mironov et al., 2010; Baldauf et
el., 2011; Wagner et al., 2016).

An online coupling between ParFlow and the COSMO model is performed via CLM3.5
(Gasper et al., 2014; Shrestha et al., 2014; Keune et al., 2016). Similar to the Common Land Model
(by (Dai et al., 2003)), CLM3.5 module accounts for surface moisture, carbon, and energy fluxes
between the shallow or near-surface soil (discretized/specification top soil layer), snow, and the
atmosphere (Oleson et al., 2008). The model components of a fully coupled system consisting of
COSMO, CLM3.5, and ParFlow are assembled by making use of the multiple–executable
approach (e.g. with OASIS3-MCT model coupler). The OASIS3-MCT coupler employs
communication strategies based on the message passing interface standards, MPI1/MPI2 and the
Project for Integrated Earth System Modeling, PRISM, Model Interface Library (PSMILe) for
parallel communication of two–dimensional arrays between OASIS3-MCT coupler and the
coupling models (Valcke et al., 2012; Valcke, 2013). The OASIS3-MCT specifies the series of
coupling, frequency of the couplings, the coupling fields, the spatial grid of the coupling fields,
transformation type of the (two–dimensional) coupled fields, and simulation time management and integration.

At the coupling interface, the OASIS3-MCT interface interchanges the atmospheric forcing terms and the surface fluxes in serial mode. The lowest level and current time step of the atmospheric state of COSMO is used as the forcing term for CLM3.5. CLM3.5 then computes and returns the surface energy and momentum fluxes, outgoing longwave radiation, and albedo to COSMO (Baldauf et al., 2011). The air temperature, wind speed, specific humidity, convective and grid-scale precipitation, pressure, incoming shortwave (direct and diffuse) and longwave radiation, and measurement height are sent from COSMO to CLM3.5. In CLM3.5, a mosaic/tilling approach may be used to represent the subgrid-scale variability of land surface characteristics, which considers a certain number of patches/tiles within a grid cell. The surface fluxes and surface state variables are first calculated for each tile and then spatially averaged over the whole grid cell (Shrestha et al., 2014). As with PF.CLM3.5, the one–dimensional soil column moisture predicted by CLM3.5 gets replaced by ParFlow’s variably saturated flow solver, so ParFlow is responsible for all calculations relating soil moisture redistribution and groundwater flow. Within the OASIS3-MCT ParFlow sends the calculated pressure and relative saturation for the coupled region soil layers to CLM3.5. The CLM3.5 also transmits depth-differentiated source and sink terms for soil moisture including soil moisture flux e.g. precipitation, and soil evapotranspiration for the coupled region soil layers to ParFlow. Applications of TerrSysMP in fully coupled mode from saturated subsurface across the ground surface into the atmosphere include a study on the impact
of groundwater on the European heat wave 2003 and the influence of anthropogenic water use on the robustness of the continental sink for atmospheric moisture content (Keune et al., 2016).

5.3 ParFlow–Weather Research Forecasting models (PF.WRF)

The Weather Research and Forecast (WRF) is a mesoscale numerical weather prediction system designed to be flexible and efficient in a massively parallel computing architecture. WRF is a widely used model that provides a common framework for idealized dynamical studies, full physics numerical weather prediction, air-quality simulations, and regional climate simulations (Michalakes et al., 1999, 2001; Skamarock et al., 2005). The model contains numerous mesoscale physics options such as microphysics parameterizations (including explicitly resolved water vapor, cloud, and precipitation processes), surface layer physics, shortwave radiation, longwave radiation, land surface, planetary boundary layer, data assimilation, and other physics and dynamics alternatives suitable for both large-eddy and global-scale simulations. Similar to COSMO, the WRF model is a fully compressible, conservative-form, non-hydrostatic atmospheric model which uses time-splitting integration techniques (discussed below) to efficiently integrate the Euler equations (Skamarock and Klemp, 2007).

The online ParFlow WRF coupling (PF.WRF) extends the WRF platform down to bedrock by including highly resolved three-dimensional groundwater and variably saturated shallow or deep vadose zone flows, and a fully integrated lateral flow above ground surface (Molders and Ruhaak, 2002; Seuffert et al., 2002; Anyah et al., 2008; Maxwell et al., 2011). The land surface model portion that links ParFlow to WRF is supplied by WRF through its land surface component, the Noah Land Surface Model (Ek et al., 2003); the standalone version of WRF has no explicit
model of subsurface flow. Energy and moisture fluxes from the land surface are transmitted
between the two models via the Noah LSM which accounts for the coupling interface, and which
is conceptually identical to the coupling in PF-COSMO. The three-dimensional variably saturated
subsurface and two-dimensional overland flow equations, and the three-dimensional atmospheric
equations given by ParFlow and WRF are simultaneously solved by the individual model solvers.

Land surface processes, such as evapotranspiration, are determined in the Noah LSM as a function
of potential evaporation and vegetation fraction. This effect is calculated with the formulation:

\[ E(x) = F f x (1 - f_{avg}) E_{pot} \]  \hspace{1cm} (22)

where \( E(x) \) stands for rate of soil evapotranspiration (length per unit time), \( f x \) represents empirical
coefficient, \( f_{avg} \) denotes vegetation fraction, and \( E_{pot} \) is potential evaporation, determined that
depends on atmospheric conditions from the WRF boundary layer parameterization (Ek et al.,
2003). The vegetation fraction is zero over bare soils (i.e. only soil evaporation), so equation 22
becomes:

\[ E(x) = F f x E_{pot} \]  \hspace{1cm} (23)

The quantity \( F \) is parameterized as follows:

\[ F = \frac{\phi s_w - \phi s_{res}}{\phi - \phi s_{res}} , \]  \hspace{1cm} (24)

where \( \phi \) is the porosity of the medium, \( s_w \) and \( s_{res} \) are relative saturation and residual saturation
respectively, from vanGenuchten relationships (VanGenuchten, 1980; Williams and Maxwell,
2011). Basically, \( F \) refers to the parameterization of the interrelationship between evaporation and
near-ground soil water content and provides one of the connections between Noah LSM and ParFlow, and thus WRF.

In the presence of a vegetation layer, plant transpiration (length per unit time) is determined as follows:

\[
T(x,z) = G(z)C_{plant}f_{veg}E_{pot},
\]

where \( C_{plant} \) represents a constant coefficient between 0 and 1, which depends on vegetation species, and the \( G(z) \) function represents soil moisture which provides other connection between the coupled models (i.e. ParFlow, Noah, and WRF). The solution procedure of PF.WRF uses an operator–splitting approach where both model components use the same time step. WRF soil moisture information including runoff, surface ponding effects, unsaturated and saturated flow, which includes an explicitly resolved water table are calculated and sent directly to the Noah LSM within WRF by ParFlow and utilized by the Noah LSM in the next time step. WRF supplies ParFlow with evapotranspiration rates and precipitation via the Noah LSM (Jiang et al., 2009). The interdependence between energy and land balance of the subsurface, ground surface, and lower atmosphere can fully be studied with this coupling approach. The coupled PF.WRF via the Noah-LSM has been used to simulate explicit water storage and precipitation within basins, to simulate surface runoffs and to simulate the land-atmosphere feedbacks and wind patterns as a results of subsurface heterogeneity (Maxwell et al., 2011; Williams and Maxwell, 2011). Studies with coupled model PF.WRF are highlighted in Table 2.

5.4 ParFlow–Advanced Regional Prediction System (PF. ARPS).
The Advanced Regional Prediction System (ARPS) composed of a parallel mesoscale atmospheric model created to explicitly predict convective storms and weather systems. The ARPS platform aids in effectively investigating the changes and predictability of storm-scale weather in both idealized and more realistic settings. The model deals with the three dimensional, fully compressible, non-hydrostatic, spatially filtered Navier-Stokes equations (Rihani et al., 2015). The governing equations include conservation of momentum, mass, water, heat or thermodynamic, turbulent kinetic energy, and the equation of state of moist air making use of a terrain-following curvilinear coordinate system (Xue et al., 2000). The governing equations presented in a coordinate system with z as the vertical coordinate are given as

\[
\frac{d\mathbf{v}}{dt} = -2\Omega \times \mathbf{v} - \frac{1}{\rho} \nabla P + g + F 
\]  

(26)

\[
\frac{d\rho}{dt} = -\rho \nabla \cdot \mathbf{v}
\]  

(27)

\[
\frac{dT}{dt} = -\frac{RT}{C_v} \nabla \cdot \mathbf{v} + \frac{Q}{C_v}
\]  

(28)

\[
P = \rho RT
\]  

(29)

Equations (26) to (29) are momentum, continuity, thermodynamic and equation of state, respectively. The material (total) derivative \(d/dt\) is defined as

\[
\frac{d}{dt} = \frac{\partial}{\partial t} + \nabla \cdot \mathbf{v}
\]  

(30)

The variables \(\mathbf{v}, \rho, T, P, g, F, Q\) in equations (26) to (29) represent velocity \([\text{LT}^{-1}]\), density \([\text{ML}^{-3}]\), temperature \([\text{K}]\), pressure \([\text{ML}^{-1}\text{T}^{-2}]\), gravity \([\text{LT}^{-2}]\), frictional force \([\text{MLT}^{-2}]\), and the
diabatic heat source [$ML^{-2}T^{-2}$], density, temperature, pressure, gravity, frictional force, and the
(diabatic heat source), respectively (Xu et al., 1991). The ARPS model employs high-order
monotonic advection technique for scalar transport and fourth-order advection for other variables
e.g. mass density and mass mixing ratio. A split-explicit time advancement scheme is utilized with
leapfrog on the large time steps, and an explicit and implicit scheme for the smaller time steps is
used to inculcate the acoustic terms in the equations (Rihani et al., 2015).

The PF. ARPS forms a fully–coupled model that simulates spatial variations in above
ground processes and feedbacks, forced by physical processes in the atmosphere and the below the
ground surface. In the online coupling process, ARPS land surface model forms the interface
between ParFlow and ARPS to transmit information (i.e. surface moisture fluxes) between the
coupled models. ParFlow as a component of the coupled model replaces the subsurface hydrology
in the ARPS land surface model. Thus, ARPS is integrated into ParFlow as a subroutine to create
a numerical overlay at the coupling interphase (specified layers of soil within the land surface
model in ARPS) with the same number of soil layers at the ground surface within ParFlow. The
solution approach employed is an operator-splitting that allows ParFlow to match the ARPS
internal timesteps. ParFlow calculates the subsurface moisture field at each timestep of a
simulation and passes the information to ARPS land surface model, which is used in each
subsequent timestep. At the beginning of each time step, the surface fluxes from ARPS that are
important to ParFlow include evapotranspiration rate and spatially–variable precipitation
(Maxwell et al., 2007). PF. ARPS has been applied to investigate the effects of soil moisture
heterogeneity on atmospheric boundary layer processes. PF.ARPS keeps a realistic soil moisture that is topographically-driven distribution and shows spatiotemporal relationship between water depth, land surface and lower atmospheric variables (Maxwell et al., 2007; Rihani et al., 2015). A summary of current studies involving PF. ARPS is included in Table 2.

5.5 ParFlow–CrunchFlow (ParCrunchFlow)

CrunchFlow is a software package developed to simulate multicomponent multidimensional reactive flow and transport in porous and/or fluid media (Steefel, 2009). Systems of chemical reactions that can be solved by the code include kinetically controlled homogenous and heterogeneous mineral dissolution reactions, equilibrium–controlled homogeneous reactions, thermodynamically controlled reactions, and biologically–mediated reactions (Steefel and Lasaga, 1994; Steefel and Yabusaki, 2000). In CrunchFlow, discretization of the governing coupled partial differential equations which connect subsurface kinetic reactions and multicomponent equilibrium, flow and solute transport is based on finite volume. (Li et al., 2007; Li et al., 2010). Coupling of reactions and transport in CrunchFlow that are available at runtimes are performed using two approaches. These are briefly discussed below.

First, a global implicit or one–step method approach is based on a backwards Euler time discretization, with a global solution of the coupled reactive transport equations using Newton’s method. This global implicit scheme solves the transport and reaction terms simultaneously (up to two-dimensional) (Kirkner and Reeves, 1988; Steefel, 2009). Second, a time or operator splitting
of the reaction and transport terms which is based on an explicit forward Euler method; the sequential non-iterative approach, SNIA (in which the transport and reaction terms are solved) (Steefel and Van Cappellen, 1990; Navarre-Sitchler et al., 2011). The stability criteria associated with the explicit approach is that the simulation timestep is restricted via the courant-Friedrichs-Lewy (CFL) condition, under the circumstance that the transportation of mass does not occur over multiple grid cell, but a single grid cell in a timestep. Thus, a small-time step must be used to ensure this condition holds. This small step size may lead to simulations that will demand much time to solve Beisman, (2007), so more processors are used, in order to decrease the processor workload and decrease solution time of the simulation. Coupling of fully saturated flow to the reactive transport calculations and coupling between a partially saturated flow and transport (flow and diffusion) can be done successively. However, these simulations require calculations of the flow and liquid saturation fields with a different model.

ParCrunchFlow is a parallel reactive transport model developed by combining ParFlow with CrunchFlow. ParCrunchFlow was designed to be only applicable for subsurface simulation. The coupled model ParCrunchFlow relies on ParFlow’s robustness ability to efficiently represent heterogeneous domains and simulate complex flow to provide a more realistic representation of the interactions between biogeochemical processes and non-uniform flow fields in the subsurface than the uncoupled model. ParFlow provides solution of Richards’ equation to ParCrunchFlow, which is not present in the biogeochemical code CrunchFlow. ParCrunchFlow employs operator-splitting method to reactive transport, in which the transport and reaction terms are decoupled and calculated independently. Online coupling between the models is achieved through a
sequential non-iterative approach, where the reaction terms in CrunchFlow’s operator-splitting solver gets connected to ParFlow’s advection terms. ParCrunchFlow takes advantage of multidimensional advection capability of ParFlow instead of CrunchFlow’s advective-dispersive transport capabilities (up to two-dimensional). A steady state governing differential equation for reaction and advection (with no dispersion and diffusion terms) in a single-phase system is given by

$$\frac{\partial c_i}{\partial t} + \nabla \cdot (\nu c_i) - R_i = 0, \quad (i = 1, N_{tot})$$

(31)

where $c_i$ is the concentration of species $i$, $\nu$ represents velocity of flow, $R_i$ indicates total reaction rate of species $i$, and $N_{tot}$ represents total species number. In the coupling process, the advection terms are calculated by ParFlow’s transport solver through a first-order explicit upwind scheme or a second-order explicit Godunov scheme. Low-order upwind weighting schemes can introduce numerical dispersion, which can impact the simulated reactions, and a comparison of several upwinding schemes can be found in (Benson et al., 2017). CrunchFlow calculates the reaction terms using the Newton-Raphson method. For example, in the coupled-model ParCrunchFlow, ParFlow code assigns all hydrological parameters, undertakes the functions relating to parallelization including domain decomposition and message transmission, and solves for pressure and flow fields. The CrunchFlow module is then used to evaluate all reaction terms and conversions between mobile and immobile concentrations. Sequence of simulations of a floodplain aquifer, comprising biologically mediated reduction of nitrate have been performed with ParCrunchFlow. The simulations demonstrate that ParCrunchFlow realistically represents the
changes in chemical concentrations seen in most field scale systems than CrunchFlow alone (summarized in Table 2) (Beisman, 2007; Beisman et al., 2015).

Figure 5 Caption: Schematic of the communication structure of the coupled models. Note: CLM represents a stand-alone Community Land Model. The modified version of CLM by Dai et al., (2003) is not shown here because it is a module only for ParFlow, not really a stand-alone LSM any longer.

6. Discussion and Summary

IHM s constitute classes of simulation tools ranging from simple lumped parameter models to comprehensive deterministic, distributed and physically based modeling systems for simulation of multiple hydrological processes (LaBolle et al., 2003; Castronova et al., 2013). They are indispensable in studying the interactions between surface and subsurface systems. IHMs that calculate surface and subsurface flow equations in a single matrix (Maxwell et al., 2015), scaling from the beginning parts to the mouth of continental river basins at high-resolutions are essential (Wood, 2009) in understanding and modeling surface-subsurface systems. IHMs have been used to address surface and subsurface science and applied questions. For example, evaluating the effects of groundwater pumping on streamflow and groundwater resources (Markstrom et al., 2008), evaluating relationship between topography and groundwater (Condon and Maxwell, 2015), coupling water flow and transport (Sudicky et al., 2008; Weill et al., 2011) and assessing
the resilience of water resources to human stressors or interventions and the variations in the 
(Maxwell et al., 2015) over large spatial extents at high resolution. Modeling or simulation at large 
spatial extents e.g. regional and continental scales and resolution e.g. 1km² (see Figure 6), and 
even small spatial scale (Figure 7) comes with the associated computational load even on 
massively parallel computing architectures. IHMs, such as ParFlow have overcome the 
computational burden of simulating or resolving questions (e.g. involving approximating variably 
saturated and overland flow equations) beyond such levels of higher spatial scales and resolutions. 
This capability may not be associated with more conceptually based models which, for example, 
may not simulate lateral groundwater flow or resolve surface and subsurface flow by specifying 
zones of groundwater network of stream before performing a simulation (Maxwell et al., 2015) 
For cross-comparison of ParFlow with other contemporary IHMs, a more comprehensive model 
testing and analyses have recently been done and readers can access these resources at Maxwell et 

Figure 6 Caption: Map of water table depth (m) over the simulation domain with two insets 
zooming into the North and South Platte River basin, headwaters to the Mississippi River. Colors 
represent depth in log scale (from 0.01 to 100 m) (Maxwell et al., 2015).

Figure 7 Caption: Map of hydraulic conductivity (K) and stream depth in the East Inlet watershed 
in Colorado (Engdahl and Maxwell, 2015). This domain covers 30km² using 3.1 million 20m² 
lateral grid cells. The springs emanating from within the hillslopes highlight the realism afforded 
by integrated modeling at small scales.
ParFlow is based on efficient parallelism (high performance efficiency) and robust hydrologic capabilities. The model solvers and numerical methods used are powerful, fast, robust, and stable, which has contributed to the code’s excellent parallel efficiency. As stated earlier, ParFlow is very capable of simulating flows under saturated and variably saturated conditions i.e. surface, vadose, and groundwater flows, even in highly heterogeneous environments. For example, in simulation of surface flows (i.e. solving the kinematic wave overland flow equations), ParFlow possess the ability to accurately solve streamflow (channelized flow) by using parameterized river routing subroutines (Maxwell and Miller, 2005; Maxwell et al., 2007, 2011). ParFlow includes coupling capabilities with a flexible coupling interface which has been utilized extensively in resolving many hydrologic problems. The interface-based and process-level coupling used by ParFlow is an example for enabling high-resolution, realistic modeling. However, based on the applications, it would be worthwhile to create one, or several, generic coupling interfaces within ParFlow to make it easier to use its surface/subsurface capabilities in other simulations. Nonetheless, ParFlow has been used in coupling studies in simulating different processes and/or systems including simulating energy and water budgets of the surface and subsurface (Rihani et al., 2010; Mikkelson et al., 2013), surface water and groundwater flows and transport (Kollet and Maxwell, 2006; Beisman, 2007; Beisman et al., 2015; Maxwell et al., 2015), and subsurface, surface, and atmospheric mass and energy balance (Maxwell and Miller, 2005; Maxwell et al., 2011; Shrestha et al., 2014; Sulis et al., 2017). Undoubtedly, such coupled-model simulations come with computational burden and ParFlow performs well in overcoming such problems, even at high
spatial scale and resolutions. This capability of ParFlow (coupling with other models) is continuously being exploited by hydrologic modelers, and new couplings are consistently being established. For example, via model coupling, the entire transpiration process could be investigated i.e. from carbon dioxide sequestration from the atmosphere by plants, subsurface moisture dynamics and impacts, to oxygen production by plants. Likewise, land cover change effects on mountain pine beetles may be investigated via coupling of integrated models. But these projected research advances can only be achieved if the scientific community keeps advancing code performance by developing, revising, updating, and rigorously testing these models’ capabilities.

Presently, ParFlow’s open source model and open developer community is fully transparent, and this openness is a major difference between it and other models that has enabled ParFlow to continue evolving. The user community is growing daily across the globe. Code developers have made available, aside from the ParFlow working manual, an active and frequently-updated blog (current blog: “http://parflow.blogspot.com/”) and other sources including “https://www.parflow.org” and “https://github.com/parflow” where code developers and experienced users provide great information and suggestions that help in fixing bugs and ease frustrations of other users. Over the years, these easily accessible resources have proven to be helpful. The code is constantly updated through release of new versions with modifications designed to meet varying hydrologic challenges and directions for applications across different scales and fields. Each ParFlow package (version) comes with verified simulation test cases with directions that simulate different real systems and idealized cases. These serve as great resource where additional code modifications have been tested in every release of the code. ParFlow has a
clear, rigorous verification procedure to make sure that any changes checked in do not “break” previous developments. This ensures numerical accuracy and backwards compatibility. Moreover, the full suite of test cases is automatically re-run before any submitted change can even be considered for merging with the master branch of the code. The number of branches/forks cannot be controlled in any open source (or community) code, but any contributions to the master branch are exhaustively vetted before being pushed out to users. Further, there is a software development and sustainability plan to improve the capabilities of ParFlow such as incorporation of new formulations of both kinematic and diffusive wave approximations, and advanced parallelization support (GPU’s and heterogeneous compute architectures). ParFlow works very well on different computing architectures and operating systems from “Laptops to Supercomputers” (single CPU, Linux clusters, highly scalable systems including IBM Blue Gene) with the same source code and input on all platforms. The code can use significant computational power and runs efficiently on supercomputing environments (e.g. Edison, Cori, JUQUEEN, and Yellowstone). Through ParFlow hydrologic modelers have available a very efficient yet still growing integrated hydrologic model to simulate and understand surface-subsurface flows.

Code availability

ParFlow is an open-source, object-oriented, parallel watershed flow model developed by community of scientists from the Environmental Protection Department at the Lawrence Livermore National Laboratory (LLNL), Colorado School of Mines and F-Z Jülich with supporting scientists from several other institutions. Versions of ParFlow are archived with detailed document or information located at: http://inside.mines.edu/~rmaxwell/maxwell_software.shtml or obtain from commercially hosted free SVN repository. Source code for the current ParFlow release “v3.5.0” can be downloaded at: https://github.com/parflow/parflow/releases/tag/v3.5.0.---The current version of ParFlow is
available at: https://github.com/parflow/parflow/releases/tag/v3.6.0. The version of ParFlow described in this manuscript is archived on zenodo: https://doi.org/10.5281/zenodo.3555297.

Author Contribution

Section 3 of the manuscript was written by Carol S. Woodward. Benjamin N. O. Kuffour and Nicholas B. Engdahl wrote the other Sections, and the entire manuscript was edited by Laura E. Condon, Stefan Kollet, and Reed M. Maxwell.

Competing Interest

We declare that no conflict of interest exist whatsoever between any of the authors and the editors or the referees.

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Figure 1: Coupled surface and subsurface flow systems. The physical system is represented on the left and a schematic of the overland flow boundary condition (continuity of pressure and flux at the ground surface) is on the right. The equation, \( p = \psi_s = \psi \) in Figure 1 signifies that at the ground surface, the vertically averaged surface pressure and subsurface pressure head are equal, which is the unique overland flow boundary used by ParFlow.
Figure 2: Representation of orthogonal (upper) and the terrain following (lower) grid formulations and schematics of the associated finite difference dependences (right). The i, j, and k are the x, y, and z cell indices.
Figure 3: Working flow chart of ParFlow’s solver for linear and non-linear system solution.
Figure 4(a): A pictorial description of the relevant physical environmental features and model coupling. CLM represents the Community Land Model, a stand-alone Land Surface Model (LSM) via which ParFlow couples’ COSMO. The modified version of CLM by Dai et al., (2003) and is not shown in Figure 4(a) because it is a module only for ParFlow, not really a stand-alone LSM any longer. The core model (ParFlow) always solves the variably saturated 3-D groundwater flow problem but the various couplings add additional capabilities.
Figure 4(b): Schematic showing information transmission at the coupling interface. PF, LSM, and ATM indicate the portions of the physical system simulated by ParFlow, Land Surface Models, and Atmospheric Models respectively. The downward and upward arrows indicate the directions of information transmission between adjacent models. Note: Coupling between ParFlow and CrunchFlow (not shown) occur within the subsurface.
Figure 5: Schematic of the communication structure of the coupled models. Note: CLM represents a stand-alone Community Land Model. The modified version of Common Land Model by Dai et al., (2003) is not shown here because it is a module only for ParFlow, not really a stand-alone LSM any longer.
Figure 6: Map of water table depth (m) over the simulation domain with two insets zooming into the North and South Platte River basin, headwaters to the Mississippi River. Colors represent depth in log scale (from 0.01 to 100 m) (reproduced from Maxwell et al., 2015). The domain uses 1km$^2$ grid cells and represents one of the largest, and highest resolution domains simulated by integrated models to date.
Figure 7: Map of hydraulic conductivity (K) and stream depth in the East Inlet watershed in Colorado (Engdahl and Maxwell, 2015). This domain covers 30km$^2$ using 3.1 million 20m$^2$ lateral grid cells. The springs emanating from within the hillslopes highlight the realism afforded by integrated modeling at small scales.
Table 1: Details for the various scaling studies conducted using ParFlow

<table>
<thead>
<tr>
<th>Simulation Case</th>
<th>Computer System</th>
<th>Processor Number</th>
<th>Jacobian/ Numerical Method</th>
<th>Preconditioner</th>
<th>Computation time (seconds)</th>
<th>Problem Size (cell Number)</th>
<th>Parallel Efficiency (%)</th>
<th>Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface processes and variably saturated flow (ParFlow and CLM)</td>
<td>JUGENE (IBM Blue-Gene Supercomputer)</td>
<td>16,384</td>
<td>Finite difference</td>
<td>ParFlow Multigrid</td>
<td>10,920</td>
<td>486,000</td>
<td>58.00</td>
<td>(Kollet et al., 2010)</td>
</tr>
<tr>
<td>Terrain Following Grid</td>
<td>JUGENE (IBM Blue-Gene Supercomputer)</td>
<td>4,096</td>
<td>Analytical</td>
<td>Non-Symmetric</td>
<td>1,130.50</td>
<td>2,048,000,000</td>
<td>80.91</td>
<td>(Maxwell, 2013)</td>
</tr>
<tr>
<td>Overland flow</td>
<td>Intel Xeon Tightly coupled Linux Cluster</td>
<td>100</td>
<td>Finite difference</td>
<td>-</td>
<td>10,800</td>
<td>50,000</td>
<td>82.00</td>
<td>(Kollet and Maxwell, 2006)</td>
</tr>
<tr>
<td>Excess infiltration produced runoff</td>
<td>Intel Xeon Tightly coupled Linux Cluster</td>
<td>100</td>
<td>Finite difference</td>
<td>-</td>
<td>10,800</td>
<td>50,000</td>
<td>72.00</td>
<td>(Kollet and Maxwell, 2006)</td>
</tr>
<tr>
<td>Terrain Following Grid</td>
<td>JUGENE (IBM Blue-Gene Supercomputer)</td>
<td>16,384</td>
<td>Finite difference</td>
<td>Symmetric</td>
<td>2,100.81</td>
<td>8,192,000,000</td>
<td>50.60</td>
<td>(Maxwell, 2013)</td>
</tr>
<tr>
<td>Subsurface and Overland flow coupling</td>
<td>IBM BGQ architecture</td>
<td>1,024</td>
<td>Analytical / Finite difference</td>
<td>ParFlow Multigrid</td>
<td>7,200</td>
<td>150,000</td>
<td>50.00</td>
<td>(Osei-Kuffuor et al., 2014)</td>
</tr>
<tr>
<td>Fully coupling terrestrial systems modeling platform</td>
<td>IBM BGQ system JUQUEEN</td>
<td>4,096</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>38,880</td>
<td>82.00</td>
<td>(Gasper et al., 2014)</td>
</tr>
<tr>
<td>Performance evaluation of ParFlow code (modified version of ParFlow)</td>
<td>(IBM Blue-Gene Supercomputer) JUQUEEN</td>
<td>458,752</td>
<td>Finite difference</td>
<td>-</td>
<td>10,569,646,080</td>
<td>-</td>
<td>-</td>
<td>(Burstedde et al., 2018)</td>
</tr>
</tbody>
</table>

Note: The hyphen "-" shows that information was not provided by the appropriate study.
### Table 2: Selected coupling studies involving application of ParFlow and atmospheric, land surface, and subsurface models

<table>
<thead>
<tr>
<th>Application</th>
<th>Coupled-Model</th>
<th>Simulation-Scale</th>
<th>Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface heterogeneity, surface energy-budget</td>
<td>CLM</td>
<td>Urban-watershed Ballona Creek watershed, CA</td>
<td>(Reyes et al., 2016)</td>
</tr>
<tr>
<td>Sensitivity analysis (evaporation parameterization)</td>
<td>CLM (modified)</td>
<td>Column</td>
<td>(Jefferson and Maxwell, 2015)</td>
</tr>
<tr>
<td>Sensitivity of photosynthesis and stomatal resistivity parameters</td>
<td>CLM (modified)</td>
<td>Column</td>
<td>(Jefferson et al., 2017)</td>
</tr>
<tr>
<td>Active subspaces; dimension reduction; energy fluxes</td>
<td>CLM (modified)</td>
<td>Hillslope</td>
<td>(Jefferson et al., 2015)</td>
</tr>
<tr>
<td>Spin-up behavior; initial conditions watershed</td>
<td>CLM</td>
<td>Regional</td>
<td>(Seck et al., 2015)</td>
</tr>
<tr>
<td>Urban processes</td>
<td>CLM</td>
<td>Regional</td>
<td>(Bhaskar et al., 2015)</td>
</tr>
<tr>
<td>Global sensitivity</td>
<td>CLM</td>
<td>watershed</td>
<td>(Srivastava et al., 2014)</td>
</tr>
<tr>
<td>Entropy production optimization and inference principles</td>
<td>CLM</td>
<td>Hillslope</td>
<td>(Kollet, 2015)</td>
</tr>
<tr>
<td>Soil moisture dynamics</td>
<td>CLM</td>
<td>Catchment</td>
<td>(Fang et al., 2015)</td>
</tr>
<tr>
<td>Dual-boundary forcing concept</td>
<td>CLM</td>
<td>Catchment</td>
<td>(Rahman et al., 2015)</td>
</tr>
<tr>
<td>Initial conditions; spin-up</td>
<td>CLM</td>
<td>Catchment; watershed</td>
<td>(Ajami et al., 2014, 2015)</td>
</tr>
<tr>
<td>Groundwater-fed irrigation impacts of natural systems; optimization water-allocation algorithm</td>
<td>CLM</td>
<td>Watershed; Sub-watershed</td>
<td>(Condon and Maxwell, 2013, 2014)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land-surface fluxes)</td>
<td>CLM</td>
<td>Watershed</td>
<td>(Condon et al., 2013)</td>
</tr>
<tr>
<td>Mountain Pine Beetle</td>
<td>CLM</td>
<td>Hillslope</td>
<td>(Mikkelsen et al., 2013)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land-surface processes)</td>
<td>CLM</td>
<td>Hillslope</td>
<td>(Atchley and Maxwell, 2011)</td>
</tr>
<tr>
<td>Computational scaling</td>
<td>CLM</td>
<td>Hillslope</td>
<td>(Kollet et al., 2010)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (infiltration in arid environment)</td>
<td>CLM</td>
<td>Hillslope</td>
<td>(Maxwell, 2010)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land-energy fluxes)</td>
<td>CLM</td>
<td>Hillslope</td>
<td>(Rihani et al., 2010)</td>
</tr>
<tr>
<td>Heat and subsurface energy transport (ParFlowE)</td>
<td>CLM</td>
<td>Column</td>
<td>(Kollet et al., 2009)</td>
</tr>
<tr>
<td>Subsurface heterogeneity on evapotranspiration</td>
<td>CLM</td>
<td>Column, Hillslope</td>
<td>(Kollet, 2009)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land-energy fluxes; runoff)</td>
<td>CLM</td>
<td>Watershed; Hillslope</td>
<td>(Kollet and Maxwell, 2008)</td>
</tr>
<tr>
<td>Climate change (land-energy feedbacks to groundwater)</td>
<td>CLM</td>
<td>Watershed</td>
<td>(Maxwell and Kollet, 2008)</td>
</tr>
<tr>
<td>Model development</td>
<td>CLM</td>
<td>Column</td>
<td>(Maxwell and Miller, 2005)</td>
</tr>
<tr>
<td>Subsurface transport</td>
<td>CLM</td>
<td>Aquifer</td>
<td>(Tompson et al., 1998, 1999; Maxwell et al., 2003)</td>
</tr>
<tr>
<td>Model development (TerrSysMP)</td>
<td>COSMO</td>
<td>Watershed</td>
<td>(Shrestha et al., 2014)</td>
</tr>
<tr>
<td>Implementation and Scaling (TerrSysMP)</td>
<td>COSMO</td>
<td>Continental</td>
<td>(Gasper et al., 2014)</td>
</tr>
<tr>
<td>Application</td>
<td>Coupled Model</td>
<td>Simulation Scale and Size (x, y, and z dimensions)</td>
<td>Model Development</td>
</tr>
<tr>
<td>----------------------------------------------------------------------------</td>
<td>---------------</td>
<td>---------------------------------------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>Groundwater response to ground surface-atmosphere feedbacks</td>
<td>COSMO</td>
<td>European CORDEX domain (Continental)</td>
<td></td>
</tr>
<tr>
<td>Atmosphere, DART, data assimilation</td>
<td>WRF</td>
<td>Watershed</td>
<td></td>
</tr>
<tr>
<td>Coupled model development (Atmosphere)</td>
<td>WRF</td>
<td>Watershed</td>
<td></td>
</tr>
<tr>
<td>Subsurface heterogeneity (runoff generation)</td>
<td>WRF</td>
<td>Hillslope</td>
<td></td>
</tr>
<tr>
<td>Subsurface uncertainty to the atmosphere</td>
<td>WRF</td>
<td>Watershed</td>
<td></td>
</tr>
<tr>
<td>Subsurface transport</td>
<td>ARPS</td>
<td>Watershed</td>
<td></td>
</tr>
<tr>
<td>Terrain and soil moisture heterogeneity on atmosphere</td>
<td>ARPS</td>
<td>Hillslope</td>
<td></td>
</tr>
<tr>
<td>Risk Assessment of CO leakage</td>
<td>CRUNCHFLOW</td>
<td>Aquifer</td>
<td></td>
</tr>
<tr>
<td>Reactive transport heterogeneous saturated subsurface environment</td>
<td>CRUNCHFLOW</td>
<td>Aquifer</td>
<td></td>
</tr>
</tbody>
</table>

Note: “CLM” show that coupling with ParFlow was by the original Common Land Model or Community Land Model. “CLM (modified)” show that the modified version of Common Land Model by (Dai et al., 2003) was a module for ParFlow.
<table>
<thead>
<tr>
<th>Groundwater-fed irrigation impacts of natural systems; optimization water allocation algorithm</th>
<th>CLM</th>
<th>Watershed; Sub-watershed (41km x 41km x 100m)</th>
<th>(Condon and Maxwell, 2013, 2014)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subsurface heterogeneity (land surface fluxes)</td>
<td>CLM</td>
<td>Watershed (209km x 268km x 3502m)</td>
<td>(Condon et al., 2013)</td>
</tr>
<tr>
<td>Mountain Pine Beetle</td>
<td>CLM</td>
<td>Hillslope (500m x 1000m x 12.5m)</td>
<td>(Mikkelson et al., 2013)</td>
</tr>
<tr>
<td>Groundwater-land surface-atmosphere feedbacks</td>
<td>CLM</td>
<td>Watershed (32km x 45km x 128m)</td>
<td>(Ferguson and Maxwell, 2010, 2011, 2012)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land surface processes)</td>
<td>CLM</td>
<td>Hillslope (250m x 250m x 4.5m)</td>
<td>(Atchley and Maxwell, 2011)</td>
</tr>
<tr>
<td>Computational scaling</td>
<td>CLM</td>
<td>Hillslope (150m x 150m x 240m)</td>
<td>(Kollet et al., 2010)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land surface fluxes)</td>
<td>CLM</td>
<td>Hillslope (32km x 45km x 128m)</td>
<td>(Maxwell, 2010)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land energy fluxes)</td>
<td>CLM</td>
<td>Hillslope (5km x 0.1km x 310m)</td>
<td>(Rihani et al., 2010)</td>
</tr>
<tr>
<td>Heat and subsurface energy transport (ParFlowE)</td>
<td>CLM</td>
<td>Column (1m x 1m x 10m)</td>
<td>(Kollet et al., 2009)</td>
</tr>
<tr>
<td>Subsurface heterogeneity on evapotranspiration</td>
<td>CLM</td>
<td>Column, Hillslope (32m x 45m x 128m)</td>
<td>(Kollet, 2009)</td>
</tr>
<tr>
<td>Subsurface heterogeneity (land-energy fluxes: runoff)</td>
<td>CLM</td>
<td>Watershed; Hillslope (3km x 3km x 30m)</td>
<td>(Maxwell and Kollet, 2008)</td>
</tr>
<tr>
<td>Climate change (land-energy feedbacks to groundwater)</td>
<td>CLM</td>
<td>Watershed (3000m x 3000m x 30m)</td>
<td>(Maxwell and Kollet, 2008)</td>
</tr>
<tr>
<td>Model development experiment</td>
<td>CLM</td>
<td>Column</td>
<td>Yes</td>
</tr>
<tr>
<td>Subsurface transport</td>
<td>CLM</td>
<td>Aquifer (30m x 15m x 0.6m)</td>
<td>(Tompson et al., 1998, 1999; Maxwell et al., 2003)</td>
</tr>
<tr>
<td>Model development (TerrSysMP)</td>
<td>COSMO</td>
<td>Watershed (64km x 64km x 30m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Implementation and Scaling (TerrSysMP)</td>
<td>COSMO</td>
<td>Continental</td>
<td>Yes</td>
</tr>
<tr>
<td>Groundwater response to ground surface-atmosphere feedbacks</td>
<td>COSMO</td>
<td>Continental (436m x 424m x 103m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Atmosphere, DART, data assimilation</td>
<td>WRF</td>
<td>Watershed (15km x 15km x 5m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Coupled model development (Atmosphere)</td>
<td>WRF</td>
<td>Watershed (15km x 15km x 5m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Subsurface heterogeneity (runoff generation)</td>
<td>WRF</td>
<td>Hillslope (3km x 3km x 30m)</td>
<td>(Meyerhoff and Maxwell, 2010)</td>
</tr>
<tr>
<td>--------------------------------------------</td>
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<td>-----------------------------</td>
</tr>
<tr>
<td>Subsurface uncertainty to the atmosphere</td>
<td>WRF</td>
<td>Watershed (15km x 15km x 5m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Subsurface transport</td>
<td>ARPS</td>
<td>Watershed (17m x 10.2m x 3.8m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Terrain and soil moisture heterogeneity on atmosphere</td>
<td>ARPS</td>
<td>Hillslope (5km x 2.5km x 80m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Risk Assessment of CO leakage</td>
<td>CRUNCHFLOW</td>
<td>Aquifer (84km x 75km x 144m)</td>
<td>Yes</td>
</tr>
<tr>
<td>Reactive transport heterogeneous saturated subsurface environment</td>
<td>CRUNCHFLOW</td>
<td>Aquifer (120m x 120m x 120m)</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Note: “CLM” show that coupling with ParFlow was by the original Common Land Model or Community Land Model. “CLM (modified)” show that the modified version of Common Land Model by (Dai et al., 2003) was a module for ParFlow.
References


Valcke, S., Balaji, V., Bentley, P., Guilyardi, E., Lawrence, B., and Pascoe, C.: Developing a


