Algorithm sensitivity analysis and parameter tuning for tissue image segmentation pipelines.

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Accessed April 15, 2020 12:19 AM EDT
Bioimage informatics

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Abstract

Motivation: Sensitivity analysis and parameter tuning are important processes in large-scale image analysis. They are very costly because the image analysis workflows are required to be executed several times to systematically correlate output variations with parameter changes or to tune parameters. An integrated solution with minimum user interaction that uses effective methodologies and high performance computing is required to scale these studies to large imaging datasets and expensive analysis workflows.

Results: The experiments with two segmentation workflows show that the proposed approach can (i) quickly identify and prune parameters that are non-influential; (ii) search a small fraction (about 100 points) of the parameter search space with billions to trillions of points and improve the quality of segmentation results (Dice and Jaccard metrics) by as much as $1.42 \times$ compared to the results from the default parameters; (iii) attain good scalability on a high performance cluster with several effective optimizations.

Conclusions: Our work demonstrates the feasibility of performing sensitivity analyses, parameter studies and auto-tuning with large datasets. The proposed framework can enable the quantification of error estimations and output variations in image segmentation pipelines.


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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Whole slide tissue images (WSIs) obtained from tissue specimens provide a means to study disease morphology at the sub-cellular scale. Several algorithm, computation and data challenges, however, have to be overcome in order to facilitate studies with large datasets of WSIs. In this work we target challenges that stem from the fact that most image analysis workflows are sensitive to variations in input parameters. A workflow optimized for a group of images may not perform well for another set of images. It is, therefore, important to (1) quantify the impact of input parameters on analysis output and (2) adjust parameters to produce more accurate analysis results. We call (1) and (2) collectively a parameter study. We refer to part (1) as the algorithm sensitivity analysis (SA) process and define it as the process of comparing results from multiple analyses of a dataset using variations of an analysis workflow (e.g. different input parameters or different algorithms) and quantifying differences in the results. Part (2) is an extension of SA and we refer to it as the parameter tuning process. We are interested in parameter auto-tuning in which the parameter space of an analysis workflow is searched automatically by generating analysis results from a set of...
Table 1. Parameters their value ranges for two example workflows

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Range value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Parameters of the Watershed based segmentation workflow</td>
<td>The parameter search space contains about 21 trillion parameter points</td>
<td></td>
</tr>
<tr>
<td>B/G/R</td>
<td>Background detection thresholds</td>
<td>[210, 220, …, 240]</td>
</tr>
<tr>
<td>TI/T2</td>
<td>Red blood cell thresholds</td>
<td>[2.5, 3.0, …, 7.5]</td>
</tr>
<tr>
<td>G1/G2</td>
<td>Thresholds to identify candidate nuclei</td>
<td>[5, 10, …, 80]</td>
</tr>
<tr>
<td>MinSize</td>
<td>Area threshold of candidate nuclei</td>
<td>[2, 4, …, 40]</td>
</tr>
<tr>
<td>MaxSize</td>
<td>Area threshold of candidate nuclei</td>
<td>[900, 1500]</td>
</tr>
<tr>
<td>MinSizePI</td>
<td>Area threshold before watershed</td>
<td>[5, 10, …, 80]</td>
</tr>
<tr>
<td>MinSizeSeg</td>
<td>Area threshold from final segmentation</td>
<td>[2, 4, …, 40]</td>
</tr>
<tr>
<td>MaxSizeSeg</td>
<td>Area threshold from final segmentation</td>
<td>[900, 1500]</td>
</tr>
<tr>
<td>FillHoles</td>
<td>propagation neighborhood</td>
<td>[4-conn, 8-conn]</td>
</tr>
<tr>
<td>MorphRecon</td>
<td>propagation neighborhood</td>
<td>[4-conn, 8-conn]</td>
</tr>
<tr>
<td>Watershed</td>
<td>propagation neighborhood</td>
<td>[4-conn, 8-conn]</td>
</tr>
<tr>
<td>(b) Parameters of the Level Set based segmentation workflow</td>
<td>The parameter search space contains about 2.8 billion parameter points</td>
<td></td>
</tr>
<tr>
<td>OTSU</td>
<td>OTSU threshold value</td>
<td>[0.3, 0.4, …, 1.3]</td>
</tr>
<tr>
<td>Curvature Weight</td>
<td>Curvature weight (CW) for level-set</td>
<td>[0.0, 0.05, …, 1.0]</td>
</tr>
<tr>
<td>MinSize</td>
<td>Minimum object size</td>
<td>[1, 2, …, 20]</td>
</tr>
<tr>
<td>MaxSize</td>
<td>Maximum object size</td>
<td>[50, 55, …, 400]</td>
</tr>
<tr>
<td>MSKernel</td>
<td>Radius in Mean-Shift calculation</td>
<td>[5, 6, …, 30]</td>
</tr>
<tr>
<td>LevelSetIter</td>
<td>Number of iterations of the level set computation</td>
<td>[5, 6, …, 150]</td>
</tr>
</tbody>
</table>

parameters, comparing the analysis results with ground truth, and repeating the process to find a set of parameters that produces the most accurate results as measured by a comparison metric.

A parameter study with a large set of WSIs can be a challenging and computationally expensive task. Consider a workflow of normalization and segmentation steps—this type of workflow is our focus in this paper, because segmentation is a crucial step in extracting salient morphology information from images and consists of several complex and parameterized data transformation operations. Processing a single WSI through this workflow can take hours on a single CPU. In addition, SA may require the evaluation of hundreds or thousands of parameter combinations because of the staggeringly high number of possible parameter values. Table 1 shows the list of parameters and their value ranges for the segmentation step in two analysis workflows used in our experimental evaluation. The first workflow uses a watershed based segmentation step (Kong et al., 2013), while the other employs level set and mean shift clustering methods (Gao et al., 2016). The large number of possible parameter values can lead to a large number of runs and high volumes of data to be handled and processed. For instance, one of our SA experiments involved 2000 runs of the watershed segmentation pipeline on 55 WSIs. The execution of the experiment on a distributed memory machine with 128 nodes took 42.9 h; and during the experiment, a total of 820 Terabytes of data were produced and processed. In studies at this scale it is imperative to utilize high performance computing systems efficiently in order to speed up the parameter study processes.

Approaches for parameter space search and parameter optimization have been developed and successfully applied in several application domains (Campolongo et al., 2007; Jones, 2001; Morris, 1991; Rios and Sahinidis, 2013; Saltelli et al., 2004; Sareni and Krähenbühl, 1998; Snoek et al., 2012; Tabatabaei et al., 2005; Weirs et al., 2012). One of the main contributions of this work is the adaptation of these methods and demonstration of their utility in the context of large-scale whole slide tissue image analysis. This is accomplished by integrating the parameter search and auto-tuning methods into a software framework that employs novel high-performance computing (HPC) techniques to address the data and computational challenges. Our contributions can be summarized as follows.

- An approach for efficient execution of sensitivity analysis for WSI segmentation workflows. Our approach implements a global SA process, which examines output sensitivity from the perspective of a range of input parameter variations (Iooss and Lemaitre, 2015). A key consideration is to reduce the number of parameter evaluations and provide relevant information about the impact of input parameters on analysis results. Our approach adapts and incorporates two types of methods (Campolongo et al., 2007; Morris, 1991; Sobol, 2001; Weirs et al., 2012) for use in WSI analysis to accomplish this: (1) methods to efficiently perform an initial and quick screening of non-influential input parameters (i.e. those parameters that do not contribute significantly to variations in output) and remove them from further consideration; and (2) methods that compute measures of importance or quantitative sensitivity indexes. The combination of these methods enables the application of sensitivity analysis with large datasets of WSIs and for segmentation workflows with large parameter spaces.

- A systematic experimental evaluation of multiple optimization algorithms for automatically tuning input parameters in segmentation workflows. Previous work on automated parameter estimation optimization in image segmentation has employed techniques for specific segmentation models (Kumar and Hebert, 2003; McIntosh and Hamarneh, 2007; Schultz and Kindlmann, 2013; Szzumener et al., 2008). The Tuner system (Torsney-Weir et al., 2011) targets general segmentation algorithms and uses statistical models from sampling runs to explore the parameter space. In our work, we treat the segmentation algorithm as a black-box and employ efficient optimization algorithms that can
quickly converge to desired results. Our results show that the output of a segmentation workflow can be significantly improved with the automated tuning approaches. In our experiments, the quality of the segmentation results was improved, as measured by the Dice (1945) and Jaccard metrics, by as much as 42% compared with the results obtained with the default workflow parameters.

- High performance computing methods to address the computation and data challenges in parameter studies. Our work provides several runtime optimizations to accelerate parameter evaluation runs on distributed-memory parallel machines. These methods are integrated in a runtime system, called Region Templates, we developed in an earlier work (Teodoro et al., 2014). The Region Templates system is designed to address the processing and data management challenges of image analysis pipelines on distributed memory systems with hybrid multi-core CPU and co-processors. The work described in this paper adds new optimizations for parameter studies: (1) efficient data movement and staging, data-aware assignment of stages and operations in an analysis workflow to optimize repeated workflow executions with different parameters; and (2) simultaneous parameter evaluation to eliminate common computations in the execution of an analysis workflow with multiple parameter sets. The combined use of these optimizations is crucial to enabling large-scale studies.

2 Methods

Our parameter study framework is illustrated in Figure 1. An investigator specifies a set of input images, an image analysis workflow, the value ranges of input parameters for the image analysis workflow, and the metric of interest (e.g. Dice) for comparison of analysis results. The image analysis workflow is executed on a high performance machine, while the input parameters are systematically varied by the framework.

The sensitivity analysis process is carried out in two phases. During these phases, multiple runs of the image analysis workflow are executed until the required set of parameters has been covered. In the first phase, a set of runs are executed and screening methods are called. The screening methods are used to determine which parameters of an analysis workflow have little impact on output variability—such parameters are called non-influential parameters (Section 2.1). The screening step is used as a filtering step with a large number of parameter values, before the more costly second phase is executed. At the end of the first phase, the investigator may remove some of the parameters from further consideration and fix their values in the second phase. The second phase computes importance measures for selected parameters (Section 2.2). This phase looks at the monotonicity and linearity of an analysis workflow’s output and correlates variance in the output with the input parameters and their first-order and higher-order effects. The results from the sensitivity analysis process are statistics that quantify variance in the analysis results as well as measures such as sensitivity indices that indicate the amount of variance in the analysis results that can be attributed to individual parameters or combinations of parameters (Saltelli, 2002; Sobol, 2001).

The parameter auto-tuning process calibrates the input parameters to generate more accurate results and requires a reference dataset (see Section 2.3). The reference dataset for tuning a segmentation pipeline can be, for example, a set of segmentation results generated by human experts. In the auto-tuning process, image analysis results (i.e. sets of segmented objects in our case) generated from a set of input parameter values are compared to the reference dataset. The comparison step computes an error estimate based on a metric, such as Dice, and feeds it to a search optimization method to generate another set of parameter values. This iterative process continues until a maximum number of iterations is reached or when the error estimate is below a threshold.

2.1 Methods to screen input parameters

Our implementation employs a commonly used screening method, called Morris One-At-A-Time (MOAT) design (Morris, 1991). This screening method perturbs each input parameter in a discretized parameter space while fixing the other input parameters. The \( k \)-dimensional input space (for \( k \) parameters) is partitioned uniformly in \( p \) levels, creating a grid with \( p^k \) points in which evaluations take place. Each perturbation of an input parameter \( x_i \) creates a parameter elementary effect (EE) computed as \( EE_i = \frac{\Delta y(x_i)}{\Delta x_i} \), where \( y(x) \) is the application output before the perturbation. In our case, the output refers to the metric of interest calculated by comparing the mask generated by the screening method to a reference mask. The reference mask is generated using the default input parameters of the image analysis workflow. To account for global SA, we use \( \Delta_i = \frac{p}{n-1} \) that leads to steps slightly larger than half of the input range for input parameters scaled between 0 and 1. The mean (\( \mu \)), modified mean (\( \mu^* \)—mean of absolute EE values) and standard deviation (\( \sigma \)) of EE are computed for each input parameter (Campolongo et al., 2007). The mean and modified mean represent the effects of an input parameter on the image analysis output, whereas the standard deviation reveals nonlinear effects. MOAT requires \( n = r\times(k+1) \) application runs (or evaluations in the parameter space), with the typical value of \( r \) being in the range of 5 to 15 (Iooss and Lemaitre, 2015).

2.2 Methods to compute importance measures

These methods calculate correlation coefficients between input parameters and application output or between pairs of input parameters. The coefficients implemented in our framework include Pearson’s correlation coefficient (CC), partial correlation coefficient (PCC), Spearman’s rank correlation coefficient (RCC) and partial
rank correlation coefficient (PRCC) (Saltelli et al., 2004). The simple and partial correlation coefficients are similar, but the latter excludes effects from other input parameters. The Spearman’s differs from the Pearson’s because the first uses ranked results. When input parameters are orthogonal, the simple and partial correlations are the same. The ranked correlations are helpful when relationships between parameters are non-linear (Saltelli et al., 2004).

The CC for \( x \) and \( y \) is calculated as:
\[
\text{Corr}(x,y) = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum (x_i - \bar{x})^2 \sum (y_i - \bar{y})^2}},
\]
where \( x \) and \( y \) are two parameters and \( \bar{x} \) and \( \bar{y} \) are their respective means. The computed value is a comparison metric value as in the screening step. Points evaluated in the input parameter space are selected from a probabilistic exploration. The framework supports the commonly used Monte Carlo sampling, Latin hypercube sampling (LHS) (McKay and Beckman, 1979), quasi-Monte Carlo sampling with Halton or Hammersley sequences, and a few other stochastic methods.

Variance-based Decomposition (VBD) sensitivity method (Weirs et al., 2012) is also available in our framework. VBD splits output uncertainty effects among individual parameters and can account for non-linear relationships among them. VBD computes the ‘main effect’ sensitivity index \( S_i \) (Sobol, 2001) and the ‘total effects’ sensitivity index \( S_T \) (Saltelli, 2002). The \( S_i \) measures the amount of variance in results that can be attributed to parameter \( i \) alone (first-order effects). If the sum of the \( S_i \) values is close to 1.0, most of the output variance is explained by single parameter effects. The total effect index \( S_T \) measures the first-order and higher-order effects due to the interaction of parameter \( i \) with the other parameters. For \( k \) input parameters and \( n \) samples, VBD requires \( n(k + 2) \) application runs, and \( n \) can be in the order of thousands (Weirs et al., 2012). It is, thus, important to remove non-influential parameters before VBD is applied.

2.3 Parameter auto-tuning algorithms

We have incorporated several parameter optimization methods in our framework and evaluated them experimentally. The methods include Nelder-Mead simplex (NM), Parallel Rank Order (PRO) (Tabatabae et al., 2005), Genetic Algorithm (GA) (Sareni and Krähenbühl, 1998), GLCCLUSTER, GLCSOLVE (Jones, 2001) and a Bayesian optimization algorithm (Snoek et al., 2012).

The NM method uses a simplex of \( k + 1 \) vertices in a \( k \)-dimensional search space. The PRO is a variant of the NM method that evaluates multiple points in the space concurrently. The Genetic Algorithm (GA) maps each parameter to a gene of an individual. The initial population is created randomly and evolved using crossover and mutation. The crossover uses a one-point crossover between pairs of individuals with a probability of \( C \). The mutation in each gene occurs with an independent probability of \( M \). The new population is evaluated and the results are input back to the GA to build another generation. The process continues until a preset number of generations (iterations) is reached. In our experiments, the probabilities \( C \) and \( M \) were empirically selected as 0.5 and 0.3, respectively, to maximize performance. We experimentally observed that adding the application default parameters to the initial population did not lead to improvements for our two use-case workflows.

The GLCCLUSTER and DIRECT methods are implemented in the TOMLAB optimization toolkit (http://tomopt.com) and provide a MATLAB interface. We implemented helper codes that wrap these methods. The helper codes are called by the parameter optimization methods with the set of parameter values (i.e. points in the parameter space) to be evaluated. The helper codes in turn invoke our framework’s runtime system to evaluate the points. A recent study has compared several optimization algorithms and showed that GLCCLUSTER obtains very good results (Rios and Sahinidis, 2013), in particular when the number of function evaluations to be performed is small. The Bayesian optimization algorithm (Snoek et al., 2012) is available from the Spearmint software (https://github.com/HIPS/Spearmint). This optimization algorithm builds and explores a probabilistic model of the function being tuned to select points in the search space to be evaluated. The search decision does not rely on local gradients or approximations only. This can benefit optimization of the parameters of complex functions. However, the cost of computing the next set of points to be evaluated can be very high with this method.

2.4 Efficient and scalable execution on parallel machines

We have implemented the sensitivity analysis and parameter auto-tuning processes as well as the analysis pipelines used in this paper in the Region Templates (RT) runtime system (Teodoro et al., 2014) for execution on HPC systems with co-processors. The RT runtime system schedules application operations across multiple computation nodes and on co-processors (such as Intel Xeon Phi). The processing structure of a region template application is expressed as a hierarchical dataflow graph. That is, an operation itself can be composed of lower-level operations organized into another dataflow. Application workflows are decomposed into operations that consume, transform and produce region template data objects (also referred to as data regions) instead of reading/writing data directly from/to other tasks or disk. In this way, the application itself does not have to deal with communication of data structures across an HPC platform. The region template data abstraction provides generic data region templates for common data structures, such as pixels, points, arrays (e.g. images or 3D volumes), segmented and annotated objects and regions, which are defined in a spatial and temporal domain. We refer the reader to our earlier publication for more details (Teodoro et al., 2014). The next sections describe two new optimizations in the RT system that address the requirements of the sensitivity study and parameter auto-tuning processes.

2.4.1 Storage management and optimizations

The hierarchical data storage (HDS) is a distributed data management infrastructure with an arbitrary number of storage levels within a node and across a distributed memory machine. The storage hierarchy is defined in a configuration file that includes the number of storage levels, the position of each level in the hierarchy and the level description: type of storage devices (e.g. RAM, SSD, HDD, etc.), capacity, etc. The HDS stores and retrieves data regions produced or requested by an application operation. Output data regions are stored automatically by the runtime system, which iterates through data regions of an operation and stores those marked as output in the highest (i.e. the fastest) level of storage with enough capacity. When a storage level is full, a cache replacement strategy selects data regions to be moved to a lower storage level. Supported data replacement policies are: First-In, First-Out (FIFO) and Least Recently Used (LRU).

We have developed a data locality-aware scheduling (DLAS) that considers the location of data when mapping analysis workflow stages to computing nodes. When an analysis operation finishes, the runtime takes into account the locality of the data produced by that operation to determine the node in which analysis operations that consume the data should be executed. DLAS calculates the amount
of data reuse by those operations and inserts the operations into a queue of preferred operations for the node where the data region was produced. A queue of preferred operations is maintained for each node in decreasing order of the amount data reuse. When a node requests work, the runtime system assigns the operation with the maximum data reuse to that node.

2.4.2 Optimized simultaneous parameter evaluation (SPE)
This optimization exploits the evaluation of multiple parameter sets in sensitivity analysis and auto-tuning studies per iteration of each process (Fig. 1). We have developed a strategy that merges and eliminates replicas of common computation paths, when multiple parameter sets are executed. Figure 2 shows two schemes for instantiating an application workflow in this scenario. The replica based scheme instantiates and executes the entire application workflow for each parameter set used. The compact composition scheme merges the instances of an application workflow into a single, compact workflow graph to reuse common steps in the separate workflow instances. Two steps are common if they use the same input data and parameter values.

The compact workflow graph representation draws from a data structure called FP-Tree (Han et al., 2000). The FP-tree represents sets of transactions in a structure in which common parts of transactions are expressed in a single path on the structure. In our case, we want to merge multiple workflows to create another workflow in which common computations from multiple parameters are eliminated. The complete algorithm to build the compact representation is provided in the Supplementary Section S2. In summary, for each parameter set to be executed, instead of creating a new independent workflow, it searches in the existing workflows for common computation paths. If a common computation path is found, it is instantiated once and its output is reused by the common path to downstream operations that will consume the data.

3 Results
We have evaluated the parameter study framework using Glioblastoma brain tumor tissue images obtained from The Cancer Genome Atlas repository for use in brain cancer studies (Kong et al., 2013). We divided the images into tiles, both because a WSI may not fit in memory and in order to distribute the processing of the WSI across multiple nodes and processors (multiple tiles are processed concurrently during the execution of an analysis workflow). We used two analysis workflows, which were made up of normalization, segmentation and comparison steps, for experimental evaluation. The first workflow employed a Watershed based segmentation step with 15 input parameters (Kong et al., 2013), while the second one used a level set method and 7 input parameters (Gao et al., 2016). Please see Table 1 for the list of parameters for each workflow. The experiments were conducted on up to 256 nodes of a distributed memory machine called Stampede (https://portal.xsede.org/tacc-stampede), which is one of the machines supported by the XSEDE consortium (https://www.xsede.org). Each node of the cluster has a dual socket Intel Xeon E5-2680 processors, an Intel Xeon Phi SE10P co-processor and 32 GB RAM. The nodes are interconnected via Mellanox FDR Infiniband switches.

3.1 Sensitivity analysis
We first employed MOAT to identify and filter out non-influential parameters, before using more costly methods, which include correlation coefficients and VBD. This first phase is referred to here as the MOAT phase. The watershed and level set based workflows processed, respectively, 55 WSIs (4276 4K × 4K image tiles) and 1 WSI (71 4K × 4K tissue image tiles) in the experiments—the tile sets did not have any background tiles with no tissue. A smaller dataset is used with the level set workflow because it uses a much more computationally expensive segmentation approach.

3.1.1 Finding important parameters with MOAT
The comparison metric in the experiments was computed as follows. An exclusive-or (XOR) operation was performed between the binary mask generated by the parameters selected by the framework and the binary mask generated by the default parameters of the workflow. The number of pixels with value 1 in the resulting binary mask was counted and used as the value of the comparison metric. We used a parameter space partition with 20 levels for each of the k parameters in Table 1, except for those that describe a propagation neighborhood and accept 2 input values only. The number of runs was calculated as \( n = r(k + 1) \), for the values of \( r \) ranging from 5 to 15. The experiments used 128 nodes of the Stampede cluster. The total execution times were 15 681s and 6825s, respectively, for the watershed and level set workflows when \( r \) was 15.

The results for the watershed workflow are presented in Table 2 in which parameters 6, 7, 8 and 14 are shown the most relevant; cause of the high values of \( \sigma \). Most of the parameters have non-linear interactions, because of the high values of \( \sigma \). Therefore, we decided to prune the list of parameters conservatively during the MOAT phase. We selected parameters T2, MaxSize, MinSizePl, MinSizeSeg in addition to G1, G2, Recon and MinSize for the more detailed and costly second phase. These parameters had at least one component \( (\mu \) or \( \sigma \) ) higher than 10\(^3\). The other ones (the rows in color red) were discarded from further analysis.

The MOAT phase of the level set workflow (shown in Supplementary Table S1 in the supplementary material because of space limitations) included a dummy parameter. This parameter was not passed to the analysis workflow as input. It was used in the experiments to quantify differences in the segmentation output due to the stochastic nature of this analysis workflow. The de-clumping phase of the level-set based segmentation approach is implemented using a randomized clustering strategy and, as a consequence, segmentation results from two runs with the same input parameter values may differ. The experimental results showed that variations in the segmentation output due to the stochastic nature of the
At Bioinformatics (were selected for further analysis, because at least one component this workflow. At the end of the MOAT phase, all the parameters other input parameters. The segmentation approach were smaller than variations due to the segmentation approach. The OTSU ratio stood out as the most significant parameter for this workflow. At the end of the MOAT phase, all the parameters were selected for further analysis, because at least one component (μ* or σ) of each parameter was above 10^8.

3.1.2 Importance measures

Pearson’s and Spearman’s Correlation Coefficients.

We computed the CC, PCC, RCC and PRCC coefficients by executing each workflow 400 times with the same dataset with different input parameter values. The runs took 27,078s and 22,696s, respectively, for the watershed and level set workflows on 128 computing nodes. The correlation coefficients between the input parameters and the analysis results are presented in Supplementary Table S2 in the supplementary material for both workflows.

For the watershed workflow, the CC values of most parameters are small (about 0.1), with the exception of OTSU (CC = 0.48). The differences between the CC and PCC values are the evidence of inter parameter correlation effects. These effects are higher for G1 and agree with the findings in the MOAT phase. The RCC values of some parameters are higher than their CC values, indicating that those parameters have a monotonic, but not a linear, correlation. This explains why some parameters assumed small CC values in the experiments. The ranking of the parameters, for instance, using RCC is similar to the ranking observed with MOAT. This confirms the MOAT results, but does not facilitate an additional parameter pruning opportunity before VBD is executed.

In the level set workflow, the OTSU was highlighted again as the most important parameter for this workflow. At the end of the MOAT phase, all the parameters were selected for further analysis, because at least one component (μ* or σ) of each parameter was above 10^8.

Variance-Based Decomposition (VBD). The Sobol’s indices are presented in Table 3 for k = 8 and k = 5 parameters (for the watershed and level set workflows, respectively) that were not filtered out in the MOAT and Correlation Coefficients phases. We used Saltelli’s approach (Saltelli, 2002) with Monte Carlo sampling. Each experiment required N = n(k + 2) runs. Because of the high computation costs, we limited the value of n to 200—this was sufficient because of small variations in the S_i indices as n increased from 100 to 200. The experiments with n = 200 and the watershed workflow required 2000 analysis runs with 55 WSIs. They took 150 890 s on 128 nodes, during which 820 TB of data were produced and consumed. The execution with the level set took 211 912 s on 128 nodes.

The results in Table 3 for the watershed workflow show that G2 has substantially more impact on output variability than the other parameters do. The sum of the S_i indices (0.74 with n = 200) is considerably smaller than 1.0, making this a non-additive model. Thus, a large fraction of the output variance cannot be attributed to a single input parameter for this workflow. The higher-order effects (S_{i_1}, - S_i) due to parameter interactions are important and cannot be ignored even if S_i is small. This is the case with Recon that has a large S_i value and a small S_i. The parameters with higher effect values (G1, G2, Recon) are used in the candidate object identification sub-step of the watershed workflow, highlighting the importance of this sub-step to the overall analysis results.

The sum of main effects is very close to 1.0 for the level set workflow; hence, the model is considered additive. The OTSU parameter explains alone most of the variability in the analysis results (S_i = 0.92 with n = 200). This parameter is also used in the candidate object identification substep in the level set workflow. The second most important parameter is the CW parameter, which is used to adapt the smoothness of the boundaries of nuclei. Our experiments show that the other parameters are less important. To validate our findings, we created a panel of segmentation results by varying the values of the two most important parameters and the value of the least important parameter for each segmentation workflow. The panel (see Supplementary Fig. S2 in the supplementary material) shows that the amount of variation in the analysis output agrees with the VBD values computed in the experiments.

In Table 4 we present a summary of the execution times of the sensitivity analysis methods. Even though the number of parameters

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### Table 2. MOAT analysis for the watershed workflow with r values of 5, 10 and 15

<table>
<thead>
<tr>
<th>Parameter</th>
<th>r = 5</th>
<th>r = 10</th>
<th>r = 15</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>μ*</td>
<td>σ</td>
<td>μ*</td>
</tr>
<tr>
<td>1-Blue</td>
<td>4.59E+04</td>
<td>1.03E+05</td>
<td>2.30E+04</td>
</tr>
<tr>
<td>2-Green</td>
<td>3.50E+04</td>
<td>7.83E+04</td>
<td>3.39E+04</td>
</tr>
<tr>
<td>3-Red</td>
<td>0.00E+00</td>
<td>0.00E+00</td>
<td>0.00E+00</td>
</tr>
<tr>
<td>4-T1</td>
<td>5.90E+05</td>
<td>9.01E+05</td>
<td>1.56E+07</td>
</tr>
<tr>
<td>5-T2</td>
<td>7.10E+07</td>
<td>1.24E+08</td>
<td>4.63E+07</td>
</tr>
<tr>
<td>6-G1</td>
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<td>5.87E+08</td>
<td>1.10E+09</td>
</tr>
<tr>
<td>7-G2</td>
<td>3.08E+09</td>
<td>3.67E+09</td>
<td>2.38E+09</td>
</tr>
<tr>
<td>8-MinSize</td>
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<td>3.24E+08</td>
<td>8.79E+08</td>
</tr>
<tr>
<td>9-MaxSize</td>
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<td>1.20E+08</td>
<td>8.18E+07</td>
</tr>
<tr>
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<td>1.32E+08</td>
<td>9.35E+07</td>
</tr>
<tr>
<td>11-MinSizeSeg</td>
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<td>1.92E+08</td>
</tr>
<tr>
<td>12-MaxSizeSeg</td>
<td>5.46E+06</td>
<td>9.33E+06</td>
<td>6.11E+06</td>
</tr>
<tr>
<td>13-FillHoles</td>
<td>5.13E+06</td>
<td>9.44E+06</td>
<td>6.95E+06</td>
</tr>
<tr>
<td>14-Recon</td>
<td>1.62E+09</td>
<td>3.12E+09</td>
<td>3.15E+09</td>
</tr>
<tr>
<td>15-Watershed</td>
<td>5.06E+07</td>
<td>5.88E+07</td>
<td>6.05E+07</td>
</tr>
</tbody>
</table>

We classify in green, yellow and red, respectively, those parameters having high, medium and low effect on the output. (Color version of this table is available at Bioinformatics online.)
is pruned through the MOAT, Correlation Coefficients and VBD phases, the execution times increase. This is because the per parameter sample size grows more rapidly than reduction in the set of parameters due to pruning. In the VBD analysis of the watershed workflow, for instance, about half of the original parameters are used, but the execution time of VBD is about 10× higher than MOAT because of the larger sample size. The execution time of MOAT could be used to estimate the execution times of the other more expensive phases by using difference in the sample sizes as a reference. This simple approach, however, tends to be less accurate when the execution time of an analysis workflow varies significantly when input parameters are changed. This is the case with the level set workflow. The ratio of sample sizes of any two phases (MOAT, Correlation Coefficients and VBD) does not reflect accurately variation in the execution time. This is less so with the watershed workflow, because the execution time of this workflow is less sensitive to the input parameters.

3.2 Parameter auto-tuning
This section evaluates the auto-tuning algorithms with the goal of maximizing the Dice and Jaccard metrics. These experiments were executed using 15 images manually segmented by a pathologist. First, we tuned the workflows for each image individually to evaluate the results generated by the automatically selected parameters. Second, we carried a random cross validation experiment, separating images into the training and test sets. In this experiment, the entire test set was analyzed using a single parameter set for each workflow. The parameters of the analysis workflows were varied within the value ranges shown in Table 1. The average values of the Dice and Jaccard metrics for all 15 images are presented in Table 5. The results for each image are provided for reference in Table S3 in the supplementary material. The auto-tuning algorithms were configured to perform a maximum of 100 function evaluations. It is also important to notice that the execution time spent by the optimization algorithm to choose the next parameter set to be evaluated varies among algorithms. It was measured to be about 77 s for the Spearman and less than 10 ms for other algorithms. Even though the difference is significant, this cost is amortized in our analysis because the application execution time to evaluate a parameter is far more expensive.

The results presented in Table 5 show that the auto-tuning algorithms significantly improved the quality of the results compared with the results generated by the default input parameters. In the watershed based workflow, the average Dice and Jaccard values were, respectively, 1.14× and 1.19× higher than the values from the default input parameters, whereas they were about 1.41× and 1.58× higher in the level set workflow. Improvements in the Dice and Jaccard values reached 22.2× and 40× depending on the image used, indicating that the results generated by the default input parameters were very poor for some images. Even though the auto-tuning algorithms improved the analysis output quality in most of the cases, an analysis of gains per image shows that no single algorithm is able to produce the best result for all the images in any of the experiments (see Supplementary Table S3 in the supplementary material). The results suggest that an ensemble of tuning algorithms should be used in order to achieve the best results instead of using a single auto-tuning algorithm. A panel with two images is presented in Figure 3 to show the improvement in segmentation output generated by the tuned versus default input parameters.

We also carried out a cross validation experiment in which we used 3 randomly selected images to train the parameter values with the GA method and the remaining 12 images to test the learned parameters. For the watershed workflow, the learned parameters improved the results compared to those from the default parameters on the testing data by 1.10× and 1.13× on average for the Dice and Jaccard metrics, respectively. The standard deviation was smaller
than 1%. For the level set based workflow, the results were even better. The average improvement amounts were 1.29/C2 and 1.42/C2. The results show that the auto-tuning algorithms could significantly improve segmentation results. In addition, the auto-tuning algorithms were able to find a good set of input parameters by examining only up to 100 parameter combinations out of billions possible.

3.3 Efficient execution of sensitivity analysis

We experimentally evaluated scalability of our framework as the configuration of hierarchical storage is varied. The experiments included storage configurations with 1 level (1L: file system—FS) and 2 levels (2L: RAM + FS) with FCFS and LRU replacement policies. We also analyzed performance of the data locality-aware coarse-grained scheduling (DLAS) as compared to using the FCFS strategy. A dataset containing 6113 4K x 4K image tiles was used with the watershed based workflow in these analyses because of space constrains. The benefits of the performance optimizations for the level set workflow are similar.

The results presented in Figure 4 show that the analysis workflow attained good scalability on a distributed memory machine. The performance of the configuration with a single storage level is faster than the ‘2L FIFO - FCFS’ due to the overhead of maintaining an extra storage level with very low data access hit rate (about 1.5%) in the RAM. However, the ‘2L - FIFO - DLAS’ configuration is better than the single level because of the higher data access hit rate (up to 72%) in the first level storage (RAM) as a result of DLAS scheduling. Moreover, the ‘2L LRU - DLAS’ resulted in the best performance with 1.17/C2 speedup on the 1L configuration due to the improved hit rate (87%). The cooperative use of CPU and Intel Phi improved performance by another 1.95/C2. The SPE optimization achieved an additional 1.6/C2 speedup.

4 Conclusions

Image analysis pipelines are sensitive to input parameters, and more effective application in research of image analysis with large datasets requires better quantification of algorithm sensitivity and tuning of image analysis parameters to produce more accurate and robust results. In this work, we demonstrate that (i) the cost of sensitivity analysis can be reduced by applying a series of optimization phases to identify and filter parameters with small impact on output variability and (ii) the cost of parameter space search can be reduced by intelligently search the space and simultaneously evaluating multiple parameter values on a cluster system while eliminating duplicate computations. We evaluate our approach on a cancer image analysis application using a large-scale cluster system. Our results show that the proposed approach can enable systematic, comparative study of analysis pipelines and improve analysis results when large datasets need to be analyzed. As a future work, we intend to integrate our framework with visual parameter optimization tools and interfaces (Pretorius et al., 2015) for a better visual analysis of results from sensitivity analysis and auto-tuning runs.

Funding

This work was supported in part by 1U24CA180924-01A1 from the NCI, R01LM011119-01 and R01LM009239 from the NLM, CNPq and NIH K25CA181503. This research used resources of the XSEDE Science Gateways program under grant TG-ASC130023.

Conflict of Interest: none declared.

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