AI-Assisted Forward Modeling of Biological Structures

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The rise of machine learning and deep learning technologies have allowed researchers to automate image classification. We describe a method that incorporates automated image classification and principal component analysis to evaluate computational models of biological structures. We use a computational model of the kinetochore to demonstrate our artificial-intelligence (AI)-assisted modeling method. The kinetochore is a large protein complex that connects chromosomes to the mitotic spindle to facilitate proper cell division. The kinetochore can be divided into two regions: the inner kinetochore, including proteins that interact with DNA; and the outer kinetochore, comprised of microtubule-binding proteins. These two kinetochore regions have been shown to have different distributions during metaphase in live budding yeast and therefore act as a test case for our forward modeling technique. We find that a simple convolutional neural net (CNN) can correctly classify fluorescent images of inner and outer kinetochore proteins and show a CNN trained on simulated, fluorescent images can detect difference in experimental images. A polymer model of the ribosomal DNA locus serves as a second test for the method. The nucleolus surrounds the ribosomal DNA locus and appears amorphous in live-cell, fluorescent microscopy experiments in budding yeast, making detection of morphological changes challenging. We show a simple CNN can detect subtle differences in simulated images of the ribosomal DNA locus, demonstrating our CNN-based classification technique can be used on a variety of biological structures.

Keywords: kinetochore, machine learning, convolutional neural network, feature extraction, forward modeling

INTRODUCTION

Forward modeling relies on the construction of an accurate simulation of a biological structure. These models are built using prior knowledge and can be compared to experimental data to either validate the model or simulate alterations to the structure. Budding yeast has proven to be a useful model of macromolecular models given the ease of fluorescent labeling of proteins and live cell imaging. Previous studies have generated models of microtubule dynamics, centromeric DNA, and the entire genome using budding yeast (Gardner et al., 2008; Wong et al., 2012; Stephens et al., 2013a,b; Lawrimore et al., 2016; Hult et al., 2017; Walker et al., 2019). However, proper assessment of a model's accuracy is complex and laborious. Here, we describe a method that incorporates computational modeling, simulated image generation, deep learning, principal component analysis, and machine learning to aid in the generation and assessment of computational models. Previous fluorescent microscopy studies utilized deep learning, reviewed...
The method described herein trains a CNN on simulated fluorescent images from computational models programmed to simulate the parameters deemed important by principal-component-analysis (PCA)-based exploratory data analysis (Figure 1) or the user. Experimental images are then classified by the CNN as a computational model. Therefore, our method utilizes CNN classification to determine which computational model best matches a given experimental image. We chose to utilize models of the budding yeast kinetochore and ribosomal rDNA locus (rDNA), but any biological structure that can be digitally imaged and spatially modeled by computer simulation can utilize this method.

The kinetochore, composed of more than 100 proteins (Biggins, 2013), is a cylindrical-shaped distribution of proteins assembled at centromeres of chromatids in eukaryotic cells and links the centromere to microtubule plus-ends (Joglekar et al., 2009; Musacchio and Desai, 2017). The kinetochore complex also plays an important role in the spindle assembly checkpoint by confirming that all the chromosomes are attached to the spindle in a bipolar orientation prior to the separation of the sister chromatids (Lara-Gonzalez et al., 2012; Joglekar, 2016; Joglekar and Kukreja, 2017). The kinetochore complex can be separated into two regions: the inner and outer kinetochore (Cheeseman et al., 2006; Fukagawa and Earnshaw, 2014; Huis in ‘t Veld et al., 2016; Musacchio and Desai, 2017). The inner kinetochore is composed of proteins that are localized close to the chromatin, interacting with the DNA. The proteins that are part of the inner kinetochore in *S. cerevisiae* include Cse4, Ame1, and Okp1 (Stoler et al., 1995; Pot et al., 2005; Hornung et al., 2014; Schmitzberger et al., 2017; Anedchenko et al., 2019). The outer kinetochore is composed of proteins that are localized close to microtubules and contain the microtubule-binding domain of the kinetochore complex. The proteins that are part of the outer kinetochore in *S. cerevisiae* include Nuf2, Ndc80, and Spc24 (He et al., 2001; Alushin and Nogales, 2011; Lampert and Westermann, 2011).

The budding yeast *S. cerevisiae* is a useful model organism for the kinetochore complex. Several of the proteins in the kinetochore complex of *S. cerevisiae* have direct homologs to other eukaryotic organisms, most importantly humans, allowing for understandings in *S. cerevisiae* to be translated to humans with relative ease (Kitagawa and Hieter, 2001). Furthermore, a large body of work on the genetics and nuclear dynamics of *S. cerevisiae* already exists, providing a useful foundation that can be further expounded upon. Lastly, while the general structure of the kinetochore complex of *S. cerevisiae* is similar to that of other eukaryotic organisms, simplifications in the structure allow for easier modeling and analysis, such as the one-to-one binding nature of a kinetochore to a microtubule that is not present in higher eukaryotic organisms (Peterson and Ris, 1976). Much work on understanding the kinetochore in the *S. cerevisiae* model has been previously performed, which include analyzing the 3D structure through electron microscopy, characterizing key kinetochore protein interactions, and identifying systematic differences between inner and outer kinetochore proteins (Peterson and Ris, 1976; Stoler et al., 1995; Pot et al., 2005; Lampert and Westermann, 2011; Haase et al., 2012; Huis in ‘t Veld et al., 2016; Joglekar and Kukreja, 2017; Musacchio and Desai, 2017).

Unlike the kinetochore, the nucleolus, the nuclear body that surrounds the ribosomal DNA locus in the budding yeast genome, appears as an amorphous crescent adjacent to the nuclear envelope in budding yeast during G1 of the cell cycle (Yang et al., 1989; Léger-Silvestre et al., 1999). Previous studies have modeled the nucleolus as a region of the genome enriched in DNA crosslinking that compacts the rDNA locus into several distinct clusters. Altering the kinetics of the crosslinks in the model changes the properties of the clusters and simulated fluorescent images of the model’s rDNA locus (Hult et al., 2017; Walker et al., 2019). Given the difficulty in detecting morphological changes in the rDNA locus/nucleolus in fluorescence images, we sought to determine if a simple CNN could correctly classify simulated fluorescence images generated from models with different DNA crosslinking kinetics.

The process of acquisition, segmentation, and analysis of microscopy images to gain insight into biological structures is limited to an inefficient and slow binary comparison of features susceptible to human bias, and does not provide any avenue for future classification of phenotypes or structures without spending large amounts of time and resources to train an individual to do so by eye. To address these concerns and provide a method that is more objective, resourceful, and time-efficient. We propose a novel pipeline (Figure 1), which is based on publicly available segmentation algorithms, deep learning and machine learning techniques, and basic statistical procedures (Versari et al., 2017).

The experimental branch of the pipeline begins by automatically detecting and segmenting budding yeast undergoing mitosis. The resulting images are then processed to remove noise and background fluorescence. An initial test on whether the physical features of the two conditions differ is then run by using a CNN. If the neural network fails to successfully both train and categorize the two conditions correctly at an accuracy of 70% or higher, then the two conditions are deemed to appear too similar for this analysis pipeline. However, if the neural network does successfully both train and categorize the two conditions, then the features that are thought to be important are extracted from the two sets of images. Principal component analysis is then used to identify the features that are of greatest importance to the distribution. These features are then validated through successful segregation of the two distributions using a support vector machine (SVM). If the SVM fails to correctly categorize the two conditions at an accuracy greater than 70% based on the features shown to be important, or if support vector machines trained on important and unimportant features have equal accuracy, then additional features are chosen and the process of extraction, identification, and validation of important features is repeated. However, if the validation is successful and only the important features build a valid separating hyperplane, those features are then used as the basis for the development of a 3D model that can output simulated microscope images. The accuracy of the 3D model is then explored through successful classification of the simulated images by a CNN that is trained on experimental images or the classification of experimental images.
FIGURE 1 | Graphical representation of AI-assisted, forward modeling method. Blue boxes denote steps in the pipeline and yellow boxes denote termination or iteration points. Green arrows depict a successful validation/classification and progression to the next step, while red arrows depict failed validation/classification and repetition of previous steps. A convolutional neural network (CNN) is a deep learning classification algorithm that learns distinguishing features directly from the images. A support vector machine (SVM) is a machine learning classification algorithm that learns from a given \( n \)-dimensional dataset of image features calculated from the images. Principal component analysis (PCA) is a statistical procedure that calculates vectors that are linear combination of the original features. The new vectors are calculated to generate the maximum amount of variance from the dataset. Each vector is ranked by the amount of variance in the transformed dataset. The importance of the original features is determined by the amount of the original feature in a vector multiplied by the amount of variance of the new vector in the transformed dataset. Italicized text on the right of the figure indicates the program associated with a given step.

by a CNN that is trained on simulated images. This procedure of building a model and analyzing it is repeated to develop and validate computational models of biological structures.

To test the pipeline's effectiveness in discerning differences in physical characteristics, we used a test case comparing the inner kinetochore, represented by Cse4, and outer kinetochore, represented by Nuf2. A simple CNN was able to distinguish experimental images of fluorescently tagged Cse4 and Nuf2. Two known metrics differentiating Nuf2 from Cse4 in mitotic yeast is the width of their distributions perpendicular to the spindle (Haase et al., 2012) and their distance from the spindle pole body (SPB), the microtubule organizing centers in yeast (Haase et al., 2013). These metrics would allow us to determine if PCA would determine these metrics as important. Indeed, these two metrics were found to be important in our PCA-based, exploratory data analysis. Lastly, we trained a CNN on simulated images of fluorescently tagged Cse4 that varied in a single model parameter, the width of Cse4 distribution. We then used the trained CNN to predict the width of the Cse4 distribution from experimental images.

To test if our method could be applied to amorphous biological structures, we tested if a CNN could distinguish simulated images from polymer models of the budding yeast nucleolus. A previous study had shown that the simulated fluorescent signal of the nucleolus increased in area, decreased in standard deviation, and exhibited fewer local maxima as the crosslinking rate increased (Walker et al., 2019), indicating the simulated, nucleolar images should be visually distinct. Indeed, our CNN was able to distinguish between simulated images created using different model parameters, demonstrating the extensibility of our method.

MATERIALS AND METHODS

Experimental Image Acquisition, Segmentation, and Normalization

Budding yeast strain YEF 473A was transformed with SPC29-RFP:HYG\textsuperscript{R} to fluorescently label the SPBs to generate the strain KBY7999. Strain KBY7999 was transformed with GFP-NUF2:NAT\textsuperscript{R} to generate strain KBY8169. Budding yeast strain YEF 473A was transformed with pKK1 to fluorescently label Cse4 with GFP and the endogenous Cse4 was removed and replaced with HYG\textsuperscript{R} to generate strain KBY2010. Strain KBY2010...
was transformed with SPC29-RFP:KAN<sup>R</sup> to fluorescently label
the SPBs and generate strain DCY1196.1. Seven Z-plane
image stacks of Spc29-RFP, N-terminal GFP-Nuf2 (KBY8169)
and Spc29-RFP, Cse4-GFP (DCY1196.1) yeast strains were
acquired with a Nikon Eclipse Ti TE2000-U inverted fluorescent
microscope using a Nikon Apo 1.4 NA 100x objective,
MetaMorph 7.8 software, Hamamatsu Orca Flash 4.0 LT
camera, and LumenCor Aura Light Engine. The cells in the
images were segmented using a MATLAB code repository,
CellStarSelect<sup>1</sup>, that utilizes the CellStar segmentation algorithm
(Versari et al., 2017) for segmenting budding yeast buds
from brightfield microscopy images. The MATLAB function
spotDetection calls the CellStar program to segment yeast buds
and uses the function advPointSourceDetection.m<sup>2</sup> (Gicconet
et al., 2017), which was based code developed for Aguet
et al., 2013), to detect kinetochore and SPB foci within
the bud segment in the fluorescent image channels. If
two kinetochore foci and two SPB foci are detected, the
bud segment is analyzed and saved in a cell array. The function compileImages.m parses the cell array containing
the segmented fluorescent images and creates a 50 × 50-
pixel image around each bud that contained two kinetochore
and SPB foci. The seven-step 50 × 50-pixel stack of each
bud was condensed into a single plane using a maximum
projection approach and had their intensity values normalized
and saved as 16-bit, RGB images. Duplicates of the images
were generated by rotating the images to generate seven
additional orientations of the initial to increase the size of the
dataset. The images then underwent a background subtraction
procedure and were de-noised using a low-pass 2D Wiener
filter. All image processing codes are contained within the
CellStarSelect repository<sup>3</sup>.

### Training and Testing of a Convolutional
Neural Network

We generated a CNN using MATLAB’s Deep Learning Toolbox
(Mathworks, Natick, MA, United States). The following
summarizes the code from the MATLAB script CNNBasic.m<sup>4</sup>. Image sets of the categories of interest were randomized and
split into training, validation, and testing data stores, with 56%
of the images used for training, 24% used for validation,
and 20% used for testing. The architecture of the CNN contained
13 layers with learn-able weights. The first 12 layers consisted
of a 4-layer pattern repeated three times, with the first layer
consisting of a 3 × 3 convolutional layer with a stride of 1 and
8, 16, and 32 filters, respectively. The second layer was a batch
normalization layer, the third layer was a rectified linear unit
layer, and the fourth layer was a 2 × 2 max-pooling layer with
a stride of 2. However, on the final repetition of the four
layers, the fourth layer was a fully connected layer, with an output size
set to the number of classification categories, that fed into a
SoftMax layer. The training used stochastic gradient descent with momentum as well as the associated default values for this

#### Feature Extraction and
Principal-Component-Analysis-Based
Feature Evaluation

Kinetochore features were extracted from the rescaled, 16-
bit, RGB, experimental and simulated images using the following
features were extracted from the images: spot height (width of
foci perpendicular to the spindle) of SPB (SPC29-RFP) foci and
kinetochore (Cse4-GFP or GFP-Nuf2), the standard deviation of
linescans of SPB and kinetochore foci parallel (X), perpendicular
(Y) to the spindle, and of a 5 × 5 cropped segment of the foci,
the mean intensity of a 5 × 5 cropped segment of the SPB and
kinetochore foci, the distance of the two kinetochore foci, the
distance of the kinetochore foci to the proximal SPB foci, and
the distance of the kinetochore foci to the proximal SPB foci
correlation.

Features of interest (Table 1) from all classification
categories were combined, normalized using MATLAB’s
built-in normalize.m function, and subjected to principal
component analysis. Principal component analysis was
performed with MATLAB’s built-in pca.m function. The
pca.m function outputted the principal components, stored in the
coeff matrix variable, and the associated percentage of the total variance explained by each principle component,
stored in the explained array variable. The importance of the
features was determined by multiplying the square of each of the principal components by the percentage of the
variance that principal component explained, and the resultant
values were ranked in descending order. These analyses were
performed by PCASimFeatureExtraction.m for comparing data
from experimental images, PCASimFeatureExtraction.m for comparing data from simulated images of inner
iketochore models, and PCASimInnerOuterFeatureExtraction.m for comparing data from simulated images of inner and outer
niketochore proteins. These functions are available at https://github.com/BloomLabYeast/FeatureExtractionAndSVM.

#### Training and Testing Binary Gaussian
Kernel

Observations and associated data from the feature extraction
were randomized and split into training and testing datasets,
with 70% of the images used for training and the remaining 30%
used for testing. The training set was then used to train a binary
Gaussian kernel-based classification model using a MATLAB
program, runSVM.m (see text footnote 5). The classifier used a support-vector-machine-based response range and a deviance
loss function with a variable regularization term strength, kernel
scale parameter, and the number of dimensions of expanded

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<sup>1</sup>https://github.com/jlaw8504/CellStarSelect

<sup>2</sup>https://github.com/HMS-IDAC/MatBots

<sup>3</sup>https://github.com/BloomLabYeast/CNNs

<sup>4</sup>https://github.com/BloomLabYeast/FeatureExtractionAndSVM
and the kinetochore microtubules. The Spc29 distribution allows the user to control several parameters of the Spc29 distribution of the Nuf2 fluorophores. The StaticKinet application allows the plus end of the microtubule. Thus, a "Length" parameter sets the range the Nuf2 fluorophores will be distributed from the plus end of the kinetochore microtubule (Cheeseman et al., 2006; Wei et al., 2017). The N-terminus of Nuf2 is bound near the minus end of the protein distributions relative to the N-terminus of Nuf2. The "Complex Diameter" parameter controls the diameter of the circular arrangement of kinetochore microtubules at their plus ends. The kinetochore microtubule plus ends can be randomly staggered by a given range, or can be staggered using a microtubule dynamics simulation using Simulink (Mathworks, Natick, MA, United States) (Stephens et al., 2013a). The "Rotation" parameter allows users to rotate the orientation of the plus ends of the kinetochore microtubules either randomly or by a fixed amount in the X, Y, and Z dimensions. The StaticKinet application was designed to simulate kinetochore protein distributions relative to the N-terminus of Nuf2. The N-terminus of Nuf2 is bound near the minus end of the kinetochore microtubule (Cheeseman et al., 2006; Wei et al., 2007; Ciferri et al., 2008). The "Length" parameter of Nuf2 sets the range the Nuf2 fluorophores will be distributed from the plus end of the microtubule. Thus, a "Length" parameter of 50 would distribute the Nuf2 fluorophores randomly in a 50 nm range from the plus end of the microtubule. The "Color Channel" parameter of Nuf2 controls the simulated color of the Nuf2 fluorophores. The StaticKinet application allows the user to control several parameters of the Spc29 distribution and the kinetochore microtubules. The Spc29 distribution allows users to approximate the structure of the spindle pole bodies (SPBs). The Spc29 parameters were designed so the user can either simulate the minus ends of the microtubules or adjust the model to create a disc-like structure by altering the structure and tubule diameters and tubule number parameters. The "Length" parameter controls the how widely the Spc29 fluorophores can distribute along the microtubule minus ends. The "Distance to (+)" parameter of Spc29 controls the average length of the kinetochore microtubules in the model, while the "Tubule Diameter" parameter of Spc29 controls the diameter of the minus ends of the kinetochore. The "Structure Diameter" parameter of Spc29 controls the diameter of the circular arrangement of the microtubule minus ends. The "Number of Fluorophore" parameter of Spc29 controls the number fluorophores per microtubule minus end. The "Number of Tubules" parameter of Spc29 controls the number of microtubule minus ends. The "Number of Tubules" parameter of Spc29 is independent of the "Number of Microtubules" parameter at the top of the interface that controls the number of microtubule plus ends. The "Color Channel" parameter of Spc29 controls the simulated color of the Spc29 fluorophores. The "Number of Arms" parameter under the kinetochore protein section controls the number of kinetochore fluorophores per microtubule plus end. The StaticKinet application models the kinetochores on a microtubule as straight rods coming from the plus ends of the microtubule to a single, defined point in space. The "Number of Bound Arms" controls the number of fluorophores simulated to be bound to a single point in space to mimic the individual kinetochore complexes converging on a single Cse4 centromere. Using the "Number of Bound Arms" parameter you can simulate a specified number of kinetochore complexes on each microtubule to randomly orient from the kinetochore plus end within a specified angle range. The length of the kinetochore fluorophore from the plus end is defined by the "Length of Arm" parameter. Users can slide the fluorophore along the simulated kinetochore "rod," toward the microtubule plus end, by adjusting

### TABLE 1 | Features of interest ranked in order of their importance by PCA.

| Feature                                      | Rank | Description                                                                 |
|----------------------------------------------|------|-----------------------------------------------------------------------------|
| Kinetochore distance to SPB                  | 1    | Euclidean distance between the brightest pixel of the kinetochore and the brightest pixel of its closest SPB foci normalized by the length of the spindle |
| Kinetochore X-distance                       | 2    | Magnitude of the distance between the brightest pixel of the kinetochore and the brightest pixel of its closest SPB foci parallel to the spindle axis, normalized by the length of the spindle |
| SPB foci mean intensity                      | 3    | Mean intensity of a 5 × 5 region encompassing the SPB foci                  |
| SPB foci height                              | 4    | Full-width-half-max of the maximum projection of the 7 × 5 region about the brightest pixel of the SPB foci perpendicular to the spindle |
| SPB foci STD in Y                            | 5    | Standard deviation of the intensities generated from a line-scan of a SPB foci perpendicular to the spindle |
| SPB foci STD in X                            | 6    | Standard deviation of the intensities generated from a line-scan of a SPB foci parallel to the spindle |
| SPB Foci STD                                 | 7    | Standard deviation of intensities of 5 × 5 region encompassing the SPB foci |
| Kinetochore foci height                      | 8    | Full-width-half-max of the maximum projection of the 7 × 5 region about the brightest pixel of the kinetochore foci perpendicular to the spindle |
| Kinetochore foci STD                         | 9    | Standard deviation of intensities of 5 × 5 region encompassing the kinetochore foci |
| Kinetochore foci STD in Y                    | 10   | Standard deviation of the intensities generated from a line-scan of a SPB foci parallel to the spindle |
| K–K distance                                 | 11   | Euclidean distance between the brightest pixels of the two kinetochore foci, normalized by the length of the spindle axis |
| Kinetochore foci mean intensity              | 12   | Mean intensity of a 5 × 5 region encompassing the kinetochore foci           |
| Kinetochore foci STD in X                    | 13   | Standard deviation of the intensities generated from a line-scan of a kinetochore foci parallel to the spindle |

3D Modeling and Simulated Imaging of the Budding Yeast Kinetochore

The StaticKinet MATLAB application (Figure 2) allows users to generated custom three-dimensional models of the budding yeast mitotic spindle and kinetochore complex. The program can model half of a mitotic spindle or an entire mitotic spindle by setting the “Number of Complexes” parameter to 1 or 2, respectively. The number and diameter of kinetochore microtubules can be set. The “Easy Align” setting forces the model to align to the X, Y, and Z axes of the visualization (bottom three panels) for easier interpretation of the model. The “Complex Diameter” parameter controls the diameter of the circular arrangement of kinetochore microtubules at their plus ends. The kinetochore microtubule plus ends can be randomly staggered by a given range, or can be staggered using a microtubule dynamics simulation using Simulink (Mathworks, Natick, MA, United States) (Stephens et al., 2013a). The “Rotation” parameter allows users to rotate the orientation of the plus ends of the kinetochore microtubules either randomly or by a fixed amount in the X, Y, and Z dimensions. The StaticKinet application was designed to simulate kinetochore protein distributions relative to the N-terminus of Nuf2. The N-terminus of Nuf2 is bound near the minus end of the kinetochore microtubule (Cheeseman et al., 2006; Wei et al., 2007; Ciferri et al., 2008). The “Length” parameter of Nuf2 sets the range the Nuf2 fluorophores will be distributed from the plus end of the microtubule. Thus, a “Length” parameter of 50 would distribute the Nuf2 fluorophores randomly in a 50 nm range from the plus end of the microtubule. The “Color Channel” parameter of Nuf2 controls the simulated color of the Nuf2 fluorophores. The StaticKinet application allows the user to control several parameters of the Spc29 distribution and the kinetochore microtubules. The Spc29 distribution allows users to approximate the structure of the spindle pole bodies (SPBs). The Spc29 parameters were designed so the user can either simulate the minus ends of the microtubules or adjust the model to create a disc-like structure by altering the structure and tubule diameters and tubule number parameters. The “Length” parameter controls the how widely the Spc29 fluorophores can distribute along the microtubule minus ends. The “Distance to (+)” parameter of Spc29 controls the average length of the kinetochore microtubules in the model, while the “Tubule Diameter” parameter of Spc29 controls the diameter of the minus ends of the kinetochore. The “Structure Diameter” parameter of Spc29 controls the diameter of the circular arrangement of the microtubule minus ends. The “Number of Fluorophore” parameter of Spc29 controls the number fluorophores per microtubule minus end. The “Number of Tubules” parameter of Spc29 controls the number of microtubule minus ends. The “Number of Tubules” parameter of Spc29 is independent of the “Number of Microtubules” parameter at the top of the interface that controls the number of microtubule plus ends. The “Color Channel” parameter of Spc29 controls the simulated color of the Spc29 fluorophores. The “Number of Arms” parameter under the kinetochore protein section controls the number of kinetochore fluorophores per microtubule plus end. The StaticKinet application models the kinetochores on a microtubule as straight rods coming from the plus ends of the microtubule to a single, defined point in space. The “Number of Bound Arms” controls the number of fluorophores simulated to be bound to a single point in space to mimic the individual kinetochore complexes converging on a single Cse4 centromere. Using the “Number of Bound Arms” parameter you can simulate a specified number of kinetochore complexes on each microtubule to randomly orient from the kinetochore plus end within a specified angle range. The length of the kinetochore fluorophore from the plus end is defined by the “Length of Arm” parameter. Users can slide the fluorophore along the simulated kinetochore “rod,” toward the microtubule plus end, by adjusting
FIGURE 2 | Kinetochore Model Graphical User Interface. The interface of the StaticKinet MATLAB application used to generate simulated fluorescent images of kine tochore models. In the bottom three panels the red dots represent fluorophores bound to the minus ends of the kine tochore microtubules (to approximate the spindle pole bodies), the blue dots represent the N-termini of Nuf2 bound to the plus ends of the kine tochore microtubule, and the green dots represent a given kine tochore protein, i.e., Cse4.

the “Point Marked” parameter. The “Angle from MT Axis” parameter controls the radial displacement of the single point in space where all the individual kine tochores converge, i.e., the Cse4 nucleosome. The parameter was meant to model the kine tochore complexes being pulled radially from the spindle. The “Help” tab allows users to access the online user guide. The “About” tab provides links to Bloom Lab code repository which contains the “KineticButShakeless” code repository that contains the StaticKinet application, the Bloom Laboratory website, and the code webpage of the Bloom Lab website. The “File” tab allows users to quit the application. The StaticKinetInMass application was used to generated multiple iterations of models designed in StaticKinet. The StaticKinetInMass program outputted a series of XML files that contained the fluorophore locations that were converted into simulated microscope images by Microscope Simulator 2 (Quammen et al., 2008) available at http://cismm.web.unc.edu/software under the “Inactive Software” section.

3D Modeling and Simulated Imaging of the Budding Yeast Genome

The budding yeast genome was simulated as a polymer bead-chain model (Hult et al., 2017; Walker et al., 2019). Each chromosome was modeled as a polymer chain, with centromeres attached to a single point on the nuclear membrane. The genome was discretized such that each bead in the model composed approximately 5 kb of DNA. The model contained 2803 beads, of which 361 comprised the rDNA locus within the nucleolus. The beads representing the ribosomal DNA locus could spontaneously crosslink with each other to mimic the crosslinking effects of the structural maintenance of chromosome proteins, i.e. cohesin and condensin. The average duration

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1https://github.com/BloomLabYeast/KineticButShakeless
2http://bloomlab.web.unc.edu/
3http://bloomlab.web.unc.edu/resources/programs/
of the crosslinks (µ) was set to either 0.09, 0.19, or 1.6 s in each simulation. The model was generated and simulated using DataTank, an object oriented programming environment8. DataTank’s microscope simulator module was used to generate simulated fluorescent images of the beads corresponding to the rDNA locus.

RESULTS

A CNN Can Distinguish Fluorescent Images of GFP-Nuf2/SPC29-RFP and Cse4-GFP/SPC29-RFP

To initially determine whether the images of inner and outer kinetochore complexes are distinguishable, images of Spc29-RFP, GFP-Nuf2 (outer kinetochore) and Spc29-RFP, Cse4-GFP (inner kinetochore) were acquired through the automated image acquisition pipeline (see section “Experimental Image Acquisition, Segmentation, and Normalization”). The automated process of acquiring, segmenting, and preprocessing images of Spc29-RFP, GFP-Nuf2 and Spc29-RFP, Cse4-GFP generated 5,920 and 4,416 images, respectively (Figure 3). The elder SPB is brighter than the younger SPB when fluorescently tagged with RFP (Pereira et al., 2001). This mismatch in SPB foci brightness causes the younger SPB foci to occasionally be erased in during the image processing (Figures 3A,B). Of the 5,920 images of Nuf2, 4,416 were randomly selected to allow for a balanced dataset. The trained CNN was able to distinguish and categorize inner kinetochore images from outer kinetochore images at a 87.4% testing accuracy (Table 2), confirming that the Nuf2 and Cse4 processed images can be distinguished by a CNN.

Ranking Kinetochore Feature Importance With Principal Component Analysis

Previous studies have shown that the foci of Cse4-GFP in metaphase yeast are, on average, further from their proximal SPB foci parallel to the spindle (Haase et al., 2013) and are “taller” (wider perpendicular to the spindle) than outer kinetochore proteins (Haase et al., 2012). Therefore, we chose to measure both the Euclidian (the one-dimensional distance that combines the distances parallel and perpendicular to the SPB foci) distance and distance parallel to the spindle of kinetochore foci to their proximal SPB and the spot height (width perpendicular to the spindle). We reasoned that the distance between the Cse4-GFP foci would be greater than GFP-Nuf2 foci given Cse4-GFP foci appear further from their proximal SPB foci; however, this metric will also change as the cell cycle progresses so we wanted to query if the feature could still distinguish Cse4-GFP and GFP-Nuf2 images. Given that foci shape has been shown to be a distinguishing criterion for fluorescently tagged inner and outer kinetochore proteins, we wanted to examine the standard deviation of signal intensity of the foci both overall and parallel and perpendicular to the spindle. Previous measurements of

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8http://www.visualdatatools.com/DataTank

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FIGURE 3 | Representative raw and processed images of GFP-Nuf2 and Cse4-GFP. (A) Raw maximum intensity projections of N-terminal GFP-tagged Nuf2 and SPC29-RFP in mitotic budding yeast cell (upper row) and raw intensity projection of CSE4-GFP and SPC29 in mitotic budding yeast cell (lower row). Arrows indicate positions of elder and younger spindle pole body foci. (B) Processed versions of maximum intensity projections in (A). Note loss of younger spindle pole body foci.
TABLE 2 | Confusion matrix of the CNN classification of processed, experimental images.

| True classification | Cse4-GFP (n = 883) | 754 | 129 |
|---------------------|--------------------|-----|-----|
| Nuf2-GFP (n = 883)  | 94                 | 789 |
| Total N = 1766      | Cse4-GFP           | Nuf2-GFP |
| Predicted classification |                |

Features all depended on the shape and intensity of the SPB foci. Statistical analysis of the processed images revealed that the GFP-Nuf2 strain’s SPB foci have higher mean intensities and have larger spot heights than SPB foci from the Cse4-GFP strain (Figures 4C–F). It is important to note that the images were rotated, rescaled, background subtracted, and de-noised using a custom MATLAB program (see section “Experimental Image Acquisition, Segmentation, and Normalization”). Therefore, the image processing enhanced differences in the appearance of SPB foci of the GFP-Nuf2 and Cse4-GFP strains. In contrast, the intensity of the kinetochore foci was one of the least important features (Table 1). This analysis was to aid in understanding what features the CNN may generate to distinguish processed images of GFP-Nuf2 and Cse4-GFP, as it is not possible to provide accurate quantitative measurements of fluorescent intensity from rescaled images.

The importance of these features was tested by training three classification algorithms using a support vector machine (SVM) with a Gaussian kernel. One set would use all the features, another would use the two most important features, kinetochore distance to SPB and kinetochore X-distance (Table 1), and another would use the two least important features, kinetochore foci mean intensity and kinetochore foci standard deviation in X (parallel to spindle) (Table 1). For each set of features, the data was randomly split 70% for training and 30% for testing, and an SVM with a Gaussian kernel trained was trained and then predicted...

FIGURE 4 | Feature comparisons between GFP-Nuf2 and Cse4-GFP processed images. Boxplots comparing kinetochore foci distance to proximal spindle pole body (SPB) foci (A), kinetochore foci distance to proximal SPB foci in X-direction only (parallel to the spindle) (B), SPB foci intensity (C), SPB foci height (width of foci perpendicular to spindle) (D), standard deviation of the intensities generated from a line-scan of a SPB foci perpendicular to the spindle (E), standard deviation of the intensities generated from a line-scan of a SPB foci parallel to the spindle (F). The three asterisks indicate a p-value < 0.001 using a Wilcoxon rank sum test.
the classifications of the test set. To deduce the ability of each feature set to distinguish images of Cse4 and Nuf2, the data was randomly split into test and training sets 30 times and an SVM was trained and tested on the resulting sets each time to generate a distribution of testing accuracies. The accuracies of the SVM’s that were trained on the two most important features were not statistically different than the accuracies of the SVM’s trained with all the features (Figure 5). In contrast, the accuracies of the SVM’s that were trained on the two least important features were statistically lower than the accuracies of the SVM’s trained with all the features (Figure 5). Thus, PCA-based, exploratory data analysis showed that distance of the kinetochore foci to the proximal SPB foci, which had been observed to be different in a previous study (Haase et al., 2013), was an important, distinguishing feature of Cse4 and Nuf2 fluorescent images. This result shows that the PCA-based method can help users identify distinguishing features of differing biological structures.

**CNN-Based Approach to Forward Modeling of the Kinetochore**

Our PCA-based, exploratory data analysis relies on feature engineering, which requires a user to generate features themselves. On the other hand, a CNN requires no feature engineering but provides little information as to why the CNN classified images into a given category. We reasoned a three-dimensional computational model that could be converted into a simulated, fluorescent image could be used to train a CNN to provide users insights into what model parameters best fit experimental images. First, a suit of differing model parameters is established and for each parameter set, multiple simulated images are created and labeled to that parameter set. Those image sets are used to train a CNN. The trained CNN then classifies experimental images to a given parameter set. Users could then examine which model parameters consistently label experimental images. To test this approach, we wanted to explore how a CNN, trained on simulated images generated by a three-dimensional, kinetochore model, would classify experimental images of Cse4-GFP. Simulated images were developed based on our current understanding of the mitotic spindle and kinetochore. We based the dimensions of the mitotic spindle on electron tomographs of the budding yeast mitotic spindle (O’Toole et al., 1999). We built our model such that the N-terminus of Nuf2 (blue) binds to the plus-end of a kinetochore microtubule, based on the reports of the N-terminus of Nuf2 binding to kinetochore microtubules (Cheeseman et al., 2006; Wei et al., 2007; Ciferri et al., 2008). In the model, Cse4 (green) is placed in line with the central axis of the microtubule by default (Figure 6A), but the exact location of Cse4 relative to the attached microtubule is unclear.
In our model, the diameter of the spindle at the kinetochore plus ends was set to 250 nm, with each microtubule having a diameter of 25 nm (Sodeik, 2000), and the microtubule ends were staggered in a uniform distribution ± 100 nm from the central line, representing the range of microtubules lengths observed in bipolar spindles (O’Toole et al., 1999). We represented the SPB (red) by placing 20 fluorophores at the minus ends of the 16 kinetochore microtubules (Figure 6A), which is a rough approximation of the complex shape of the yeast SPB (Bullitt et al., 1997; Muller et al., 2005; Viswanath et al., 2017). Our computational model is converted to a fluorescent image using Microscope Simulator 2 (Quammen et al., 2008), which converts each Cse4 (green) and SPB (red) bead in Figure 6A into a single fluorophore positioned in three dimensions (Figure 6B) and then into a simulated, fluorescent image (Figure 6C) by convolving each fluorophore with a point-spread function. The spreading of the light via diffraction blurs the distributions of the fluorophores into individual foci (Figure 6C).

**TABLE 3** | Confusion matrix of the CNN classification of simulated Cse4 images of varying radial displacements.

| True classification | Sim. 0 nm/On-axis (n = 14,987) | Sim. 25 nm (n = 14,832) | Sim. 50 nm (n = 14,706) | Sim. 100 nm (n = 14,130) |
|---------------------|-------------------------------|-----------------------|------------------------|------------------------|
| Sim. 0 nm/On-axis   | 14,984                        | 0                     | 0                      | 0                      |
| Sim. 25 nm          | 0                             | 14,825                | 7                      | 0                      |
| Sim. 50 nm          | 0                             | 2                     | 14,703                 | 1                      |
| Sim. 100 nm         | 0                             | 0                     | 0                      | 14,130                 |
| Total N = 58,655    |                               |                       |                        |                        |

**TABLE 4** | Categorization of experimentally acquired inner kinetochore images by the CNN trained on simulated images of the inner kinetochore of varying radial displacements.

| Exp. Cse4 | Sim. 0 nm/On-axis | Sim. 25 nm | Sim. 50 nm | Sim. 100 nm | Predicted classification |
|-----------|-------------------|------------|------------|-------------|-------------------------|
| Total N   | 4,416             | 14,984     | 14,832     | 14,706      | 14,130                  |

**FIGURE 7** | Kinetochore model organization with Cse4 radially displaced from kinetochore microtubule axis. (A–C) Scatter plots depicting the end-on view of the 25, 50, and 100 nm radial displacement models. Cse4s are the green dots, N-termini of Nuf2 are the blue dots forming a ring around the kMT plus-ends, and the SPBs are composed of red dots surrounding the minus-ends of the kMTs. (D–F) Volumetric models, generated by the Microscope Simulator 2 program, of the 25, 50, and 100 nm radial displacement organizations that were used for simulated, fluorescent image generation. (G–I) The respective simulated microscope images to the volumetric models shown in panels (E,F). White bars indicate Cse4 foci height.
A controversial parameter in our model is the position and number of Cse4 molecules. The initial assumption in the field was that there was one Cse4-containing nucleosome (2 Cse4 proteins per nucleosome) per centromere in budding yeast (Meluh et al., 1998; Collins et al., 2004; Joglekar et al., 2008b). Using independent fluorescent standards, subsequent studies found the signal of metaphase Cse4 foci was brighter than expected (Coffman et al., 2011; Lawrimore et al., 2011) and that Cse4 foci were “taller” (wider perpendicular to the spindle) than outer kinetochore foci (Haase et al., 2012). These findings suggest that Cse4 proteins are spread perpendicular to the spindle in budding yeast during metaphase. Therefore, we altered the placement of Cse4 to gradually increase its radial distance from the corresponding kinetochore microtubule center in our computational model (Figures 7A–C). The radial models were converted into Microscope Simulator 2 simulations (Figures 7D–F) and then into simulated, fluorescent images (Figures 7G–I). The increase in radial displacement of Cse4 did result in noticeably “taller” Cse4 foci in the simulated images (Figures 7D–F). We trained a CNN on simulated images generated from models were Cse4 was displaced by 0 nm (Figure 6), 25 nm, 50 nm, or 100 nm (Figure 7). The validation accuracy of the CNN on simulated images was 99.98%, demonstrating the CNN could distinguish the change in Cse4 radial position with high accuracy (Table 3). We used the trained CNN to predict the radial displacement of 4,416 processed experimental images of Cse4-GFP and SPC29-RFP (Table 4). Most processed images were classified with a radial displacement of 25 or 50 nm, with only 1.9% (83 of 4,416) of images being classified as 0 nm. This result demonstrates that our CNN-based approach to forward modeling captured the biological observation that Cse4-GFP foci are spread perpendicular to the spindle axis. Thus, this CNN-based approach to forward modeling is suitable for fine tuning computational models of biological structures.

**CNN Classification of Simulated Nucleolar Images**

The nucleolus is the region within the nucleus that surrounds the rDNA locus. Fluorescently labeled nucleolar proteins appear amorphous, making identification of morphological phenotypes difficult. Recently, two studies utilized three-dimensional, computational simulations of the budding yeast genome to study how different DNA crosslinking kinetics within the rDNA locus affect nucleolar sequestration and rDNA structure (Hult et al., 2017; Walker et al., 2019). We reasoned our CNN-based, forward modeling approach would be appropriate for this computational model if a CNN could detect the subtle differences in the simulated images generated from models with different crosslinking kinetics (Figure 8). We found that the CNN was able to identify the average crosslinking duration with 99.76% accuracy (Table 5). Thus, this rDNA model and CNN architecture would be suitable for CNN-based forward modeling.

**DISCUSSION**

**The Kinetochore as a Test-Case for AI-Assisted, Forward Modeling**

Here we demonstrate a method that combines deep learning and computational modeling to compare models of biological structures directly with experimental images (Figure 1). A relatively simple CNN, built with MATLAB’s Deep Learning ToolBox (Mathworks, Natick, MA, United States), was able to determine the difference between GFP-Nu2/Spc29-RFP image and Cse4-GFP/Spc29-RFP images with 87.4% accuracy (Table 2). To validate the difference between the images of GFP-Nu2/Spc29-RFP and Cse4-GFP/Spc29-RFP, we extracted features that were known to be different between the two strains. The importance of these features was determined using PCA and validated by comparing the classification accuracies of

![Image](image-url)
SVM’s trained on features deemed important versus SVM’s trained on features deemed unimportant. Indeed, the SVM’s trained on unimportant feature performed worse than those trained on important features (Figure 5). The most important feature was the distance of the kinetochore foci to the proximal SPB foci (Table 1), that had previously been shown to be different between Cse4 and Nuf2 (Haase et al., 2013). Thus, this PCA-based exploratory data analysis can detect biologically relevant difference between fluorescent images. However, the PCA-based analysis can only indicate a feature as discernable. The PCA-based analysis cannot discern the appropriate value of a given feature. In order to determine the measurements of features we generated a three-dimensional, computational model of the kinetochore. We then altered the radial displacement of Cse4 in the model to determine which radial displacement settings fit the experimental images. The CNN classified 98.1% of experimental images of Cse4-GFP in the 25 nm and 50 nm radial displacement categories (Table 4). This result recapitulated the biological finding that Cse4-GFP is “taller” (wider perpendicular to the spindle) than other fluorescently labeled outer kinetochore proteins (Haase et al., 2012).

The kinetochore model illustrated that a CNN used in conjunction with a three-dimensional model can be used to fine-tune model parameters to match experimental images. We used the kinetochore as test case for our AI-assisted forward modeling pipeline, but any biological structure that can be imaged and computationally modeled can be analyzed by this method. Unlike the kinetochore, images of fluorescently labeled nucleolar proteins often appear amorphous, making CNN-based forward modeling particularly useful. We again found that a relatively simple CNN, built with MATLAB’s Deep Learning ToolBox (Mathworks, Natick, MA, United States), was able to determine the difference between simulated nucleolar images (Figure 8) generated from a budding yeast genome model (Hult et al., 2017; Walker et al., 2019) with 99.76% accuracy (Table 5). Thus, a CNN is capable of discerning differences in amorphous structures as well.

Future Directions for AI-Assisted Forward Modeling

Here, we only altered a single parameter, the radial displacement of Cse4, in our computational model. However, it is possible to alter multiple parameters of our model and train a CNN on the different combinations allowing us to fine tune the model on multiple parameters. An alternative approach is to use a system similar to a generative adversarial network (GAN) (Goodfellow et al., 2014). GANs have been used recently to generate synthetic super-resolution images from lower resolution microscopy images (Ouyang et al., 2018; Wang et al., 2019; Zhang et al., 2019). In this approach the computational model would be driven by a generative algorithm to generate simulated images. A discriminative algorithm that had been trained on experimental images would then classify the simulated image into a given class or flag it as a detected fake. Since each simulated image is derived from a model, the generative algorithm could automatically tune model parameters to best fit an experimental image dataset. The generated images could then be interrogated by feature extraction to ensure key feature metrics were maintained in the simulated image generation. This GAN-based approach is a natural extension of the method we have described here and could be used on any biological structure that can be imaged and computationally modeled.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

JL provided technical guidance for the computer simulations and construction of the analysis pipeline, implemented the analysis, and wrote the manuscript. AD built the kinetochore simulation and analysis pipeline, generated the implemented analysis, and wrote the manuscript. BW constructed and ran the budding yeast genome simulations and generated the simulated images of fluorescently labeled rDNA loci. KB and JL provided the biological framework and assisted with the completion of the manuscript.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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