

Methods

High-throughput measurement of plant fitness traits with an object detection method using Faster R-CNN

Peipei Wang^{1,2}* (D), Fanrui Meng^{1,2}* (D), Paityn Donaldson¹, Sarah Horan¹, Nicholas L. Panchy³ (D), Elyse Vischulis⁴, Eamon Winship⁵ (D), Jeffrey K. Conner^{1,6,7} (D), Patrick J. Krysan⁸ (D), Shin-Han Shiu^{1,2,4,7,9} (D) and Melissa D. Lehti-Shiu¹ (D)

¹Department of Plant Biology, Michigan State University, East Lansing, MI 48824, USA; ²DOE Great Lake Bioenergy Research Center, Michigan State University, East Lansing, MI 48824, USA; ³National Institute for Mathematical and Biological Synthesis, University of Tennessee, 1122 Volunteer Blvd, Suite 106, Knoxville, TN 37996-3410, USA; ⁴Genetics and Genome Sciences Graduate Program, Michigan State University, East Lansing, MI 48824, USA; ⁵Department of Biochemistry and Molecular Biology, Michigan State University, East Lansing, MI 48824, USA; ⁶W.K. Kellogg Biological Station, Michigan State University, 3700 E. Gull Lake Drive, Hickory Corners, MI 49060, USA; ⁷Ecology, Evolution, and Behavior Graduate Program, Michigan State University, East Lansing, MI 48824, USA; ⁸Department of Horticulture, University of Wisconsin-Madison, Madison, WI 53705, USA; ⁹Department of Computational Mathematics, Science, and Engineering, Michigan State University, East Lansing, MI 48824, USA

Summary

Authors for correspondence: Shin-Han Shiu Email: shius@msu.edu

Melissa D. Lehti-Shiu Email: lehtishi@msu.edu

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• Revealing the contributions of genes to plant phenotype is frequently challenging because loss-of-function effects may be subtle or masked by varying degrees of genetic redundancy. Such effects can potentially be detected by measuring plant fitness, which reflects the cumulative effects of genetic changes over the lifetime of a plant. However, fitness is challenging to measure accurately, particularly in species with high fecundity and relatively small propagule sizes such as *Arabidopsis thaliana*.

• An image segmentation-based method using the software IMAGEJ and an object detectionbased method using the Faster Region-based Convolutional Neural Network (R-CNN) algorithm were used for measuring two Arabidopsis fitness traits: seed and fruit counts.

• The segmentation-based method was error-prone (correlation between true and predicted seed counts, $r^2 = 0.849$) because seeds touching each other were undercounted. By contrast, the object detection-based algorithm yielded near perfect seed counts ($r^2 = 0.9996$) and highly accurate fruit counts ($r^2 = 0.980$). Comparing seed counts for wild-type and 12 mutant lines revealed fitness effects for three genes; fruit counts revealed the same effects for two genes.

• Our study provides analysis pipelines and models to facilitate the investigation of Arabidopsis fitness traits and demonstrates the importance of examining fitness traits when studying gene functions.

Introduction

A major goal of biology is to understand the molecular basis for the development of organisms and their adaptation to different environments (McDonald, 1983). One approach is to evaluate the effects of genetic variants on phenotypes. However, it is often challenging to investigate such effects because gene functions may be masked by genetic redundancy (Bouché & Bouchez, 2001; Sun *et al.*, 2012) and/or be condition specific (Hirsch *et al.*, 1998; Meissner *et al.*, 1999). Moreover, the physiological and/or developmental changes caused by loss of gene

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function may be too subtle to detect. This challenge can be alleviated by measuring the effects of genetic variations on fitness (i.e. the ability of an individual to survive and reproduce) because it reflects the cumulative effects of genetic changes over the lifetime of a plant. Accurate estimates of fitness are therefore valuable for several fields of study, including plant genetics, evolution and plant breeding.

Among fitness measures, the most direct measure is the number of progenies produced (Thomson & Hadfield, 2017). In *Arabidopsis thaliana*, a predominantly selfing plant, the total number of seeds produced per plant is a particularly good estimate of fitness because it incorporates both male and female contributions. However, because Arabidopsis seeds are small (c. 0.1–0.2 mm²;

^{*}Joint first authors.

Jahnke *et al.*, 2016) and produced in large numbers (up to thousands per plant; Boyes *et al.*, 2001; Morales *et al.*, 2020), it is difficult to obtain accurate seed counts. As a consequence, fruit (silique) number (Busoms *et al.*, 2015) and total fruit length (Roux *et al.*, 2004; Busoms *et al.*, 2015; Kerwin *et al.*, 2015) are often used to measure fitness. Both measures are correlated with seed production, but fruit number is not perfectly correlated with seed number (e.g. $r^2 = 0.960$, Mauricio & Rausher, 1997) and correlations with fruit length are highly variable across studies, ranging from $r^2 = 0.988$ (Roux *et al.*, 2004) to $r^2 = 0.256$ (Gnan *et al.*, 2014). In addition, fruit numbers (up to 450 per plant; Hamidinekoo *et al.*, 2020) are typically counted manually, and these counts can be error-prone. Thus, to better measure fitness, both fruit and seed numbers should be evaluated using methods that are not hindered by propagule size or number.

Several programmes have been designed to increase the efficiency and accuracy of seed analyses. Some are aimed at measuring the properties of individual seeds (e.g. size and shape) and others at obtaining high-throughput seed counts (Herridge et al., 2011; Tanabata et al., 2012; Moore et al., 2013). These approaches typically require that seeds be separated before imaging, which increases the time needed for processing. Other systems have been designed to separate seeds mechanically such as the phenoSeeder device (Jahnke et al., 2016), large-particle flow cytometer (Morales et al., 2020) and the BELT imaging system combined with the phenoSEED algorithm (Halcro et al., 2020). A drawback of these methods is that they require specialized equipment, hindering their widespread adoption. Another approach that has been increasingly used in plant biology for applications such as measurement of fitness traits is machine vision, the application of deep learning algorithms to image analvsis (Mochida et al., 2019).

Deep learning approaches, in particular Convolutional Neural Network (CNN)-based frameworks, have been developed to detect vastly different objects (from cars to plant seeds) in images. For example, aiming to train instance segmentation models where seed counting was not the primary task, Toda et al. (2020) were able to detect the seeds of rice, lettuce, oat and wheat with 96% recall and 95% precision using Mask Region-based CNN (R-CNN). However, the detection of much smaller objects using CNN-based approaches remains challenging (Cao et al., 2019), likely because CNNs create low-level abstractions of the images, and if the objects are too small, the resulting abstractions are too simple to be used to distinguish whether the object is present or not. Although the CNN-based models developed by Toda et al. (2020) detected seeds with high accuracy, the smallest seeds tested were lettuce seeds, which have areas ranging from 1.5 to 3.6 mm² (Penaloza et al., 2005) and are c. 10 times larger than Arabidopsis seeds. Another consideration is that the most convenient way to count all the seeds from an Arabidopsis plant, which can produce thousands of seeds (Boyes et al., 2001; Morales et al., 2020), would be to put all the seeds in a single image, thus resulting in a relatively small ratio of seed size to image size. However, because of the small images $(1024 \times 1024 \text{ or } 2000 \times 2000 \text{ px}^2)$ used in Toda et al. (2020), the ratio of seed size to image size was relatively large (> 5000 px^2 per barley seed), which limited the

number of seeds that could be included in an image. Therefore, it is important to assess how well the CNN-based approaches perform in detecting objects as small as Arabidopsis seeds in an image containing thousands of them.

Convolutional Neural Network-based approaches have also been used in fruit counting. For example, wheat spikes can be detected, counted and analysed to estimate yield using R-CNN (correlation between true and predicted counts: $r^2 = 0.93$ with a slope of 1.01; Hasan et al., 2018). Starting from two pretrained models (ResNet and ResNext), Afonso et al. (2020) applied the Mask R-CNN approach to detect and count tomato fruits from images, obtaining an F1 of 0.94 when fruits partially overlapped with each other. DeepPod effectively counts Arabidopsis fruits but results in a high number of false negatives when there are many fruits ($r^2 = 0.90$ with a slope of c. 0.70; Hamidinekoo et al., 2020). In addition, the inflorescences need to be harvested when the fruits are still green, preventing the harvesting of seeds for future propagation or analysis. Thus, it is important to develop tools or models to detect and count mature fruits when seeds need to be saved for future experiments. Because Arabidopsis fruits shatter easily when dry, such tools should ideally be able to count fruits at different stages, including intact fruits and those that have already dehisced and released seed.

In this study, we evaluated two approaches for counting seeds from an Arabidopsis plant in a single image: (1) a segmentationbased method using the software IMAGEJ (Schneider *et al.*, 2012) and (2) an object detection method using the Faster R-CNN algorithm (Ren *et al.*, 2017). We also applied Faster R-CNN to count fruits in images of whole plants captured after seeds were mature. To facilitate seed and fruit counting in diverse images, we established models using input images with varying resolution, contrast, brightness and blurriness. The final seed and fruit models are provided and can be readily used by the research community. Finally, we used our pipeline to count seeds for loss-offunction mutants of six pairs of duplicate genes. We showed that mutation of three genes affects fitness, illustrating the potential importance of measuring fitness traits and the utility of our pipeline in the investigation of gene functions.

Materials and Methods

Plant materials

T-DNA insertion mutants in the Arabidopsis Col-0 background and wild-type (WT) Col-0 controls were used for training seed and fruit counting models. Information about these lines is provided in Supporting Information Tables S1–S3. Fitness data are reported for T-DNA insertion mutants of *PURPLE ACID PHOSPHATASE* 2 (*PAP2*), *PAP9*, *HIGH MOBILITY GROUP A4* (*HON4*, also known as *GH1-HMGA1*), *HON5* (*GH1-HMGA2*), *EUKARYOTIC INITIATION FACTOR 4B1* (*EIF4B1*), *EIF4B2*, *ADENOSINE 5 -PHOSPHOSULFATE REDUCTASE-LIKE 5* (*APRL5*), *APRL7*, *PLANT AND FUNGI ATYPICAL DUAL-SPECIFICITY PHOSPHATASE 3* (*PFA-DSP3*), *PFA-DSP5*, *KINESIN 7.2* (*KIN7.2*) and *KIN7.4* (Tables S4, S5). These mutants were collected as part of a large-scale study to assess the degree of genetic redundancy between duplicate genes. Multiple homozygous mutant and WT sibling plants were identified by PCR with gene-specific primers (two to six plants per genotype, Table S3). Seeds harvested from these independent lines (referred to as sublines) were planted (n = 5-20 per subline, total $n \ge 40$ per genotype) for fitness comparison between mutants and WT, and each mutant was compared with its WT sibling. This was done to reduce the chance that observed fitness effects were due to other undetected T-DNA insertions.

For plants grown for fitness analysis (Tables S3–S5) and seed scan images (Table S1), seeds were grown as described in Methods S1. Plants were grown until they were mature (i.e. had undergone global arrest). When plants were completely dry, the numbers of intact and completely or partially shattered fruits from each plant were recorded as detailed in Methods S1. The total number of seeds produced per plant was estimated in two steps. First, the number of seeds counted was divided by the number of intact fruits to obtain the average seed number per fruit. Second, the average seed number per fruit was multiplied by the total fruit number (both intact and shattered) to estimate the total seed number per plant. Plants used for fruit imaging (Table S2) were grown as described in Methods S1.

Seed image scanning, processing and counting with the segmentation method

Before seed imaging, we separated the seeds from the chaff (see Methods S1). Seed images were obtained by placing Petri plate lids containing seeds in a template made from white acrylic (295 mm \times 210 mm \times 10 mm, Fig. 1a) and taking scans with a desktop scanner (see Methods S1). The IMAGEJ (v.1.52a, https://imagej.nih.gov, Schneider *et al.*, 2012) workflow is shown in Fig. 1. Details about seed counting using IMAGEJ are provided in Methods S1. The image conversion programme and the IMAGEJ macro were combined into a Windows batch script (available in our GitHub repository, see the 'Data availability' section), in which a for-loop was used to quickly count seeds for images in sequence. It took *c*. 5 min to fully process 10 images.

Seed image processing and counting with an object detection method using Faster R-CNN

Before seed detection, each scanned image was split into 12 subimages; each subimage contains a single plate lid and is



Fig. 1 Workflow and performance for seed counting with a segmentation method using IMAGEJ when seeds were deliberately separated. (a) Workflow. Seeds from 12 different plants were scattered and manually separated from each other on the lids of 12 Petri plates, which were placed in a template and scanned. Twelve search areas, each with a diameter of 60 mm (yellow circles), were predefined. A threshold was applied by selecting pixels with intensities between 50 and 140 to separate the seed areas (red) from the background. Pixels were then converted to real-world distance units in mm. The 'Analyze Particles' tool was used to detect and count the seeds. (b) An example of an image with detected seeds (left) and an enlarged image showing the seeds (right). Red region with number, individual detected seed area. (c) Correlation between true and predicted seed counts using the segmentation method when seeds were deliberately separated.

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referred to as a 'whole-plate image'. After testing several algorithms, we chose to use Faster R-CNN for seed detection (for reasons, see Methods S1). Faster R-CNN combines the generation of region proposals (i.e. circumscribing the areas of interest, a regression problem) and their classification (i.e. in our case, the object is a seed or not) into a single pipeline (Ren *et al.*, 2017). In Faster R-CNN, images were first processed by a feature extractor (INCEPTION v.2; Szegedy *et al.*, 2016), and the resulting feature maps were used to predict bounding boxes (referred to as proposals) containing images of individual seeds (left panel in Fig. S1); these proposals were then used to crop features from the feature maps (right panel in Fig. S1). These cropped features were subsequently used for classification and bounding box regression.

Faster R-CNN models were trained using TENSORFLOW object detection API (Huang *et al.*, 2017) and implemented in TENSOR-FLOW v1.13.2 (Abadi *et al.*, 2016) in PYTHON v3.6.4. In the initial Faster R-CNN modelling trial, each whole-plate image was split into four quarter-plate images. Images were preprocessed, and seeds were annotated as detailed in Methods S1. To speed up the training process, a pretrained model (faster_rcnn_inception_v2_coco, https://docs.openvino.ai/latest/omz_models_model_faster_ rcnn_inception_v2_coco.html) was used as a starting point. To optimize Arabidopsis seed detection, we conducted hyperparameter tuning (Methods S1; Tables S6, S7; Figs S2, S3) and evaluated tuned models using the measure IoU, which is defined as the intersection (I) over (o) the union (U) of a ground truth area and a prediction area, as detailed in Methods S1.

Fruit image capturing and counting with an object detection method based on Faster R-CNN

Each dry Arabidopsis plant was placed on a pink paper background and photographed with an iPhone 8 smartphone. The images were saved in jpeg format with dimensions of 3024×4032 pixels. Fruits in the images were manually annotated, and the annotated coordinates were then converted to the csv and TFrecord formats, as conducted for the seed images (Methods S1). The same pretrained Faster R-CNN model used for seed counting was used to build the fruit counting models, and the same three hyperparameters were tuned to optimize the model performance but with a different hyperparameter space (Table S8). For each hyperparameter combination, a model was saved after 6000 steps, when the performance had converged. A final model was established using hyperparameters selected based on performance on the validation set images.

Statistical analysis of fitness traits

Data from the border cells (see Methods S1) showed different distributions compared with data from inside cells; therefore, these data were excluded from further analysis. For each block (i.e. one including *pap*, *hon* and *eif4b* and one including *aprl*, *pfa-dsp* and *kin7*, see Methods S1), quantile normalization was performed across flats using the R package 'BROMAN' (https://github.com/kbroman/broman) to account for variation between flats. Each mutant was compared with its WT control using the

Wilcoxon rank-sum test. Each pair of duplicate genes had the same WT sibling control.

Results

Seed counting with the segmentation method using IMAGEJ

Because IMAGEJ is widely used for seed morphology analysis (Cervantes *et al.*, 2016), we first developed a pipeline for seed counting that incorporated IMAGEJ analysis based on segmentation of seed areas. When fewer than 200 seeds were placed on the plate lid and separated using forceps, seeds were detected and counted with high accuracy (correlation between true and predicted seed counts, $r^2 = 0.998$, slope = 0.9998, 60 images, Fig. 1b,c; Table S9). Our segmentation-based pipeline allowed the detection of *c*. 52 template images (total of 624 plate lids) per hour with a typical laptop (Intel Core i7-7500 U CPU, 16GB RAM).

However, when seeds were placed on plate lids without separation, big clumps of seeds were not counted by the segmentation method, and small clumps where a small number of seeds were touching each other were recognized as single seeds (Fig. 2a). The prediction accuracy drops off as the number of seeds increases (Fig. 2c; Table S10); this is because the more seeds there are on the plate lid, the more likely it is that seeds touch each other, leading to an increase in the false-negative rate of prediction. Moreover, the detection of seeds could be disrupted by scratches or letters on the plate lids, and seeds outside the predefined circular search regions were not detected (purple arrowheads in Fig. S4). Thus, to obtain accurate counts based on segmentation, it is necessary to separate seeds and confine them to the centre of the plate lid, which is time-consuming and not amenable to high-throughput analysis.

Improved seed counting by an object detection method based on Faster R-CNN

We then evaluated the performance of an object detection approach using Faster R-CNN in seed counting. Since it is timeconsuming to annotate a large number of seeds for model training, we adopted a two-step strategy. First, we split the 256 whole-plate images into 1024 quarter-plate images and manually labelled a subset (180) of these quarter-plate images to speed up the training process. A total of 160 labelled quarter-plate images (*Training image set 1* in Fig. 3a) were used to build the models, and the remaining 20 images were set aside as the *Validation image set* (Fig. 3a) to evaluate model performance. A model (Model_{seed} 66) built with the optimal hyperparameter combination (scale-B, aspect ratio-A and 10 000 proposals, see Methods S1) was used to detect seeds in the remaining 844 quarter-plate images to produce *in silico* seed annotations for the second-round modelling (Fig. 3a, b), resulting in 211 labelled whole-plate images.

A new model, Model_{seed} 67, with the same parameters as $Model_{seed}$ 66, was built using 161 (*Training image set 2* in Fig. 3b) out of these 211 images. The remaining 50 labelled whole-plate images (*Test image set* in Fig. 3b) were used to evaluate the performance of Model_{seed} 67, which had an improved

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Fig. 2 Comparison between the performances of the segmentation and Faster R-CNN-based seed counting methods for test set images of seeds that were not deliberately separated. (a, b) The same seed scan image analysed by the segmentation method using IMAGEJ (a) and by Faster R-CNN (b). Three different regions of the plate lid with different densities are outlined. Region 1 has low seed density, region 2 has moderate density and region 3 has a high density. In (a), the red coloured regions represent the segmented areas identified by the segmentation method; seeds outlined in yellow and assigned numeric IDs were counted. In (b), the blue rectangles represent seeds detected by Faster R-CNN. (c, d) Correlation between true and predicted seed numbers from segmentation method (c) and Faster R-CNN (d) analysis of the test set. (e) Distribution of differences between true and predicted seed numbers. Red line, the segmentation method using IMAGEJ; blue line, Faster R-CNN. (f) Correlation between seed density index (SDI) and difference between true and predicted seed counts. Each dot in (c, d, f) corresponds to one of the 50 test set images. The red line in (c) is the regression line obtained using the loess method. The blue lines in (d, f) are fitted regression lines for Faster R-CNN predictions. The red line in (f) is the fitted linear regression line for the segmentation method-based predictions. PCC, Pearson correlation coefficient.

average F1 of 0.992 (Table S10) compared with the F1 (c. 0.970) of Model_{seed} 66 (Fig. S2). Note that the test set images were not used for training or validating Model_{seed} 67; they were thus ideal for independently testing the model. In contrast to the segmentation method, Model_{seed} 67 correctly predicted seeds even if they were in contact with each other (Fig. 2b), and the prediction accuracy was not influenced by the total seed number ($r^2 = 0.9996$, P = 1.7e-83, Fig. 2d). The differences between true and predicted seed counts were close to zero, much smaller than those in segmentation-based analysis (Fig. 2e). Furthermore, Model_{seed} 67 allowed the detection and counting of seeds in c. 240 whole-plate images per hour using 1 GPU (NVIDIA Tesla

K80) with 4 GB of GPU memory in a UNIX cluster, or *c*. 33 images per hour using a laptop with 16 GB of memory (i.e. *c*. 800 seed images can be processed per day). These results suggest that our Faster R-CNN-based models provide highly accurate Arabidopsis seed counts and can be used for large-scale fitness studies.

Impact of seed density on the Faster R-CNN model

The number of seeds in an image has a detrimental effect on the performance of the segmentation method, but not on that of Faster R-CNN (Fig. 2d). To verify that the Faster R-CNN model



Fig. 3 Workflow for building Faster R-CNN-based seed counting models. (a) First-round modelling for increasing the number of annotated seed labels. Each of the 256 whole-plate images was split into four quarter-plate images. Among the 1024 quarter-plate images, 180 were used in first-round modelling, and the remainder (844) were used in second-round modelling described in (b). Seeds in the 180 quarter-plate images were manually annotated, and these annotated images were further split into training set 1 (160) and a validation set (20) to train and evaluate models, respectively. Sixty-three combinations of three hyperparameters (i.e. 3 proposal numbers \times 3 scales (A, B and C) \times 7 aspect ratios (AR-A through G); for scale and aspect ratio values, see Supporting Information Table S6) were used to build 63 models. The optimal scale (B) and aspect ratio (AR-A) were selected based on the model performance on validation set images (Fig. S2). An additional three models (Model_{seed} 64–66) were built using scale B, AR-A and three larger proposal values, and the final best model, Model_{seed} 66, with 10 000 proposals, was applied to the 844 quarter-plate images reserved for second-round modelling to generate *in silico* seed annotations. (b) Second-round modelling. The 844 quarter-plate images with seed predictions from Model_{seed} 66 were rejoined together to reconstruct 211 using 161 (training set 2) out of the 211 annotated images with the same hyperparameters used in Model_{seed} 66, and was evaluated using the test set (i.e. the 50 independent timages plus 1700 images modified from the test set images that had different image properties (blurriness, brightness, contrast and resolution values)). For data augmentation, the image properties of 20 images from training set 2 were modified, and the resulting 420 images were combined with training set 2 (161 images), resulting in 581 images (modified training set 2), which were used to build Model_{seed} 68. The modified test set was used to evaluate the performance o

performance was not affected by the seed density, we established the seed density index (SDI), which takes into account the differing densities across a single plate. First, a circle with a radius of 30 pixels (corresponding to 0.62 mm, approximate length of two seeds) was drawn from the centre of a seed, and then, the number of seeds with central points located within the circle was calculated. Finally, the average number of seeds per circle in a wholeplate image was defined as the SDI (Fig. 4a).

We calculated the SDIs of the test set images (e.g. see Fig. S5) and determined the Pearson's correlation coefficient (PCC) between SDI and the performance of Modelseed 67 on the test set images (Fig. 4b). The higher the seed density, the lower the model performance (PCC between SDI and F1 was -0.581, P = 9.8e-06, Fig. 4b; for the correlation between SDI and other performance measures, see Fig. S6). Nevertheless, the effect of seed density on the performance of Model_{seed} 67 was small, as the F1 only dropped from 1.000 for an SDI of 1.157 to 0.971 for an SDI of 3.100 (Fig. 4b; Table S10). An F1 of 0.971 with a recall of 0.968 indicates that for an image with 1000 seeds, there would only be 32 false negatives (seeds not detected) and 25 false positives (seeds detected in an area with no seeds or a seed area counted more than once). Consistent with this, there was no significant correlation between the SDI and the difference between true and predicted seed counts (PCC = -0.206, P = 0.15), in contrast to the significant negative correlation observed for the segmentation method (PCC = -0.886, P = 1.2e-17, Fig. 2f). We also calculated SDIs for the predicted seed coordinates and found that the PCC value between true and prediction-based SDIs was 0.997 (P=1.5e-54, Fig. 4c), demonstrating that our Faster R-CNN model also predicts the locations of seeds very well.



Fig. 4 Effect of seed density on the performance of the Faster R-CNN models. (a) Examples with different seed density index (SDI) values. The radius of each circle is 30 pixels (0.62 mm). (b, c) Relationship between SDI and model performance (b) and between the true SDI and SDI based on prediction (c) for test images. Each dot corresponds to one of 50 test set images. Blue lines are the fitted linear regression lines. F1, F1 value at 0.5 IoU (intersection over union). PCC, Pearson correlation coefficient.

Model improvement through data augmentation

Our goal is to provide a seed counting model that can be widely used by different researchers, who may have seed images with different properties. Thus, we investigated the utility of Model_{seed} 67 using images with varying resolution, contrast, brightness and blurriness (Fig. 5a). These modified seed images were created by modifying the properties of the test set images (Fig. 3b, for the image property settings, see Table S11). In the modified test set, there were 1750 images: the original test set images (50) and modified images with 34 different attributes $(34 \times 50, \text{ light})$ green box, Fig. 3b). A slight but significant decrease in F1 was observed when the brightness of the images was ≤ 0.60 (*P* = 0.01, one-sided Wilcoxon signed-rank test) relative to the original images, while the F1 dropped dramatically when the relative brightness was ≥ 1.20 (P = 6.4e-08, Fig. 5b). A significant decrease in F1 was also observed when the relative contrast of images (relative to the original image) was ≤ 0.50 (*P*=1.0e-07) or ≥ 1.75 (P=5.0e-4), the relative blurriness was ≥ 1.50 (P=6.7e-10) or the relative resolution was ≤ 0.50 (P=9.1e-10), Fig. 5b). These results suggest that although Modelseed 67 is suitable for a range of image qualities, the seed detection accuracy will decrease dramatically when the image properties deviate from the training images beyond a certain point.

To improve the robustness of Model_{seed} 67, we applied data augmentation, a method used to increase the size of a training data set by including images with more properties so that better prediction models can be built (Shorten & Khoshgoftaar, 2019). To accomplish this, we used 20 of the 161 training set 2 images to produce additional images with 21 different property settings $(21 \times 20$, darker green box, Fig. 3b; for the image property settings, see Table S11). These 420 additional images, together with the original 161 images, were used to build a new model, Model_{seed} 68 (Fig. 3b), with the same hyperparameter settings as Model_{seed} 67. Model_{seed} 68 was then used to detect seeds in the modified test set images. Although there was a slight decrease in F1 when the relative blurriness was ≥ 3.00 (*P*=0.04, median F1 decrease = 0.002) or when the relative resolution was ≤ 0.30 $(P=0.02, \text{ median F1 decrease} = 0.003, \text{ Fig. 5b}), \text{ Model}_{\text{seed}} 68$ (blue, Fig. 5b) performed better than the non-augmented Modelseed 67 (red, Fig. 5b) in all situations, and thus, the augmented model is robust to different image properties.

Fruit counting using Faster R-CNN models

Compared with seed number, total fruit count is an even more frequently used proxy for fitness. Because dry Arabidopsis fruits shatter easily, it is not always possible to harvest all fruits produced by a single plant after seeds have matured, especially for plants growing in the field. In this case, the best method would be to count all fruits (including dehisced ones) and count seeds per fruit for a subset that have not dehisced, and then calculate total seed number by multiplying the number of seeds per fruit by the total fruit number. Thus, to obtain more accurate estimates of seed production per plant, it is necessary to record the numbers of both intact and shattered fruits. With these



Fig. 5 Improvement of model robustness using training images with different properties. (a) Examples of seed images with different relative brightness, contrast, blurriness and resolution values that were derived from the same original image. (b) Model performance for Model_{seed} 67 and Model_{seed} 68 on the modified test set (Fig. 2b). F1, F1 value at 0.5 IoU (intersection over union); red boxplot, Model_{seed} 67; blue boxplot, Model_{seed} 68; horizontal line in the box, median value; box range, interquartile range (IQR), that is 25^{th} (Q1) to 75^{th} percentile (Q3); whisker below box, Q1–1.5×IQR to Q1; whisker above box, Q3 to Q3+1.5×IQR.

considerations in mind, we developed Faster R-CNN models to count all fruits without harvesting the fruits first. When capturing the images for fruit counting, a pink background was used to maximize the contrast between the background and the dark, dry fruits and the pale replum of shattered fruits that remained after the valves fell from the fruit (Fig. 6a,d). Because fruits in each image were less abundant and much larger compared with seeds, we manually labelled the fruits in 120 images.

Eighty, 20 and 20 images were randomly selected and used as training, validation and test sets, respectively (Fig. 6a). Different combinations of hyperparameter values (Table S8) were evaluated, and the resulting models (Model_{fruit} 1-75, Fig. 6a) had similar performances with an average F1 of 0.925 (Fig. S7). Thus, to minimize the computational cost (lower scales or aspect ratios) while maximizing the number of fruits detected per plant (more proposals), the model built with scale_{fruit}-A, aspect ratio_{fruit}-A and 500 proposals (Model_{fruit} 21) was used. Model_{fruit} 21 was applied to the test set images, resulting in an average F1 of 0.914 (Table S12). This F1 value translates into one false positive and 15 false negatives for an image with 100 fruits. Although the r^2 between true and predicted fruit counts was 0.980 (P = 6.7e-17), the detection error increased with an increasing number of fruits in an image and the error was mostly due to undercounting or false negatives (Fig. 6b,c). The majority of the false negatives were unopened fruits that overlapped with the stem or with each other. One potential reason for the failure to detect these fruits is that they are similar to the stem in colour and shape. Another reason may be the smaller number of labelled intact fruits (543)

compared with the number of pale replums (2082) in our training images.

To assess the robustness of our model on images with different qualities, we applied Model_{fruit} 21 on test set images with different image properties (Fig. 6d). In this modified test set, there are 700 images: the original test set images (20) and modified images with 34 different attributes (34 \times 20, for the image property settings, see Table S11). Significant decreases in F1 were observed when the relative image brightness was ≤ 0.70 (P = 0.04) or ≥ 1.40 (P = 0.02), the relative contrast was $\leq 0.50 \ (P = 0.02)$ or $\geq 1.50 \ (P = 0.03)$, the relative blurriness was ≥ 2.0 (P=0.002) or the relative resolution was ≤ 0.6 (P = 0.05) (Fig. 6e). By including images with different properties (Table S11) in the training set (1840 images), a new model, Modelfruit 76, was established and applied to the modified test set. A significant but slight decrease in the resulting F1 values was only observed when the relative resolution was ≤ 0.3 (*P* = 0.02, median F1 decrease = 0.01) (Fig. 6e), indicating the robustness of Model_{fruit} 76. Using this model, 180 images could be processed per hour using a UNIX node with 1 GPU and 4 GB graphics memory, and 90 images per hour could be processed using a laptop (1 CPU, 16 GB memory). Thus, our Faster R-CNN-based models can process over a thousand plant images per day.

Effects of loss of gene function revealed by measuring fitness traits

To evaluate the importance of fitness traits in investigating gene functions and the utility of our pipeline, the fruits and

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Fig. 6 Fruit counting using Faster R-CNN models. (a) Fruit counting workflow. (b) Relationship between true and predicted fruit numbers. (c) Relationship between fruit number in an image and the model performance. PCC, Pearson correlation coefficient. (d) Examples of the same fruit image with different relative brightness, contrast, blurriness and resolution values. (e) Model performance for $Model_{fruit} 21$ and $Model_{fruit} 76$ on test images with different properties. F1, F1 at 0.5 IoU (intersection over union); red boxplot, $Model_{fruit} 21$; blue boxplot, $Model_{fruit} 76$; horizontal line in the box, median value; box range, interquartile range (IQR), that is 25^{th} (Q1) to 75^{th} percentile (Q3); whisker below box, $Q1-1.5 \times IQR$ to Q1; whisker above box, Q3 to Q3+1.5 $\times IQR$.

seeds produced by loss-of-function mutants of six pairs of duplicate genes (Tables S3–S5) were counted and compared with those of WT. Of these 12 mutants, three (*pap2, kin7.4* and *hon5*) showed a significant difference in total seed count compared with the corresponding WT control (Figs 7, S8, S9). One of these genes, *PAP2*, modulates carbon metabolism; in addition, overexpression of *PAP2* resulted in earlier bolting and a higher seed yield than WT (Sun *et al.*, 2012), which is consistent with the lower fitness that we observed for the *pap2* mutant (total seed counts, P = 3.6e-03, Wilcoxon rank-sum test, Fig. 7b). However, when studying this same mutant, Sun *et al.* observed

no significant differences in plant growth or seed yield relative to WT (Sun *et al.*, 2012).

One possible explanation for this discrepancy is the different fitness measures used by Sun *et al.* (2012) – seed weight per plant, weight per 100 seeds and fruit number per plant – none of which were significantly different between *pap2* and WT in their study. To compare our fitness estimates more directly with those of Sun *et al.*, we measured the same traits and found no significant difference in fruit number (P= 0.15, Fig. 7a) or total seed weight per plant (P= 0.40, Fig. 7c). However, the *pap2* mutant did have a higher weight per 100 seeds than the WT (P= 3.8e-08,



Fig. 7d). This could potentially indicate differences in viability because larger seeds have more resources for germination and early seedling growth (Sundaresan, 2005), but we observed no difference in germination rate between WT and *pap2* (Table S4), suggesting that there is no difference in seed viability. Taken together, our findings suggest that seed number is a better measure for revealing fitness effects of loss of *PAP2* function. However, we cannot rule out the possibility that we observed these effects because our experimental conditions were more stressful (i.e. nutrient limiting) than those in Sun *et al.* (2012).

For HON5, which encodes a high-mobility group protein (Kotliński et al., 2017), and KIN7.4, which belongs to the kinesin motor family, members of which are involved in microtubule-based movement (Moschou et al., 2016), there were significant differences in both fruit numbers (P = 0.04 for hon5 and P=5.0e-04 for kin7.4, Fig. 7e,g) and seed numbers (P=5.8e-03 for hon5 and P=3.0e-05 for kin7.4, Fig. 7f,h) between the mutants and WT. No functions have been reported for KIN7.4. HON5 was previously shown to regulate the transition to flowering along with HON4 by repressing FLC expression, but no effects on fitness were reported (Zhao et al., 2021). Loss of function of HON4 was previously reported to cause sterility (Charbonnel et al., 2018), but neither we (Fig. S8g-i) nor Zhao et al. (2021) observed this phenotype when using a different mutant with an insertion in a similar location (intron 2), suggesting that the sterility phenotype of the hon4 mutant may be dependent on environmental conditions.

Discussion

Fitness is one of the best measures of gene functionality because it reflects the ability of a plant to survive and reproduce given all Fig. 7 Fitness measurements for three mutants. (a-d) Fruit counts per plant (a), seed counts per plant (b), seed weight per plant (c) and weight per 100 seeds (d) for the T-DNA insertion mutant of PURPLE ACID PHOSPHATASE 2 (pap2) and wild-type (WT). (e, f) Fruit (e) and seed (f) counts per plant for the T-DNA insertion mutant of HIGH MOBILITY GROUP A5 (hon5) and WT. (g, h) Fruit (g) and seed (h) counts per plant for the T-DNA insertion mutant of KINESIN 7.4 (kin7.4) and WT. Sample sizes are shown in parentheses on the x-axis. P-values are from Wilcoxon signed-rank tests. Horizontal line in the box. median value; box range, interquartile range (IQR), that is 25th (Q1) to 75th percentile (Q3); whisker below box, $Q1-1.5 \times IQR$ to Q1; whisker above box, Q3 to Q3+1.5 \times IQR; violin plot, distribution of datapoint values; dot, datapoint from an individual plant; yellow, loss-of-function mutant; cyan, WT.

the phenotypic effects of the mutation over the lifetime of the individual. For self-pollinating species such as Arabidopsis, fitness is better assessed by counting the numbers of seeds than fruits, as they more directly reflect the number of offspring and reproductive success. Because of the lack of an effective tool enabling high-throughput counting of small seeds en masse, seed counts are often estimated indirectly, for example, by dividing the total seed weight per plant by the estimated individual seed weight (Cvetkovic et al., 2017) or multiplying the fruit count by the average fruit length (Kerwin et al., 2015; Taylor et al., 2019). However, these approaches may not yield accurate estimates of seed production because of potential variation in seed size, such as that between *pap2* and WT (Fig. 7c,d), and the imperfect correlation between seed number and fruit length (Roux et al., 2004). Here, we established a model employing a deep learning approach, Faster R-CNN, to count Arabidopsis seeds - one of the smallest objects analysed using machine vision to date - with a near perfect accuracy (F1 = 0.992) using images with multiple different properties or qualities.

Our model outperforms the Mask R-CNN approaches in Toda *et al.* (2020) (F1 of *c.* 0.95), where the detected objects were much larger than Arabidopsis seeds. Mask R-CNN is built on top of Faster R-CNN, so the differences in performance likely are not due to differences in algorithms. The better performance of our model is likely because our training seed images are more representative of the diversity in seed sizes and shapes than the repetitive cropped images used by Toda *et al.* The Faster R-CNN-based predictions greatly outperform those of the segmentation method implemented in IMAGEJ, a well-known platform with macros/modules for segmentation and morphology extraction (Schneider *et al.*, 2012; Cervantes *et al.*, 2016; Vasseur *et al.*, 2018). In addition, object detection based on Faster R-CNN is

less time-consuming than segmentation using IMAGEJ because seeds can be accurately detected without first being separated or confined to predefined regions.

One of the challenges when using deep learning approaches is the requirement for a large number of labelled data (in our case, labelled seeds). To overcome this, we adopted a two-step modelling strategy to reduce the labour needed for seed annotations. In step 1, we split the images and used a subset of the split images to build a preliminary model (F1 < 0.975) and applied it to the remaining images. While the predictions were not perfect, this step drastically reduced the manual annotations needed because we only needed to correct mis-predictions to boost our seed labels by *c*. 5-fold (29 360 labels in the first round, 138 929 labels in the second round). Using this much larger set of seed labels, new models were built (step 2) that had improved model performance (F1 = 0.992), indicating the effectiveness of our strategy.

The Faster R-CNN approach also shows promise in fruit detection and counting ($r^2 = 0.98$, slope = 0.79). The performance of our fruit counting model was better than that of another recently published CNN-based approach, DeepPod ($r^2 = 0.90$, slope c. 0.70, Hamidinekoo et al., 2020). In that paper, the task (i.e. fruit detection) was first divided into four classification tasks: the detection of the tip, body and base of the fruits and the detection of the stem. The separately detected parts were then joined together as a whole fruit. As the authors noted, this post-processing step affected the final fruit detection performance. In our study, the fruits were labelled and detected as whole objects, thus avoiding the need for post-processing. In addition, different from Hamidinekoo et al. (2020), where most of the fruits and stems in the images were fresh and green, fruits in our study were dry and light brown to grey, or were shattered with only the pale replum remaining. Thus, our fruit counting approach is expected to be applicable to a wider range of Arabidopsis fruit developmental stages. This is especially important when plants must be grown to maturity, and seed counts are estimated by multiplying the average number of seeds per intact fruit by the total number of fruits (intact and dehisced) (Conner & Rush, 1997).

Nevertheless, our fruit counting models did not perform as well as our seed counting models and a published IMAGEJ-based segmentation and skeletonization approach ($r^2 = 0.91$, slope ≈ 1 ; Vasseur et al., 2018), which may be due to the many fewer labelled fruits than labelled seeds (there were c. 52 times more labelled seeds than fruits). Thus, the performance of the fruit counting model is expected to be improved when more fruit labels are included to train the model. In addition, one notable drawback of our approach is the undercounting at higher fruit numbers; this was mainly due to overlap between intact fruits and between intact fruits and stems. To remedy this, one approach is to rearrange the inflorescences before capturing the images to keep fruits from overlapping with each other and with stems. Another potential approach, which is an important future direction, is to analyse multiple images (or frames of a movie) taken at different angles or to examine the 3D reconstruction of the inflorescence. In addition, there have been substantial advances in object detection algorithms in terms of performance and processing speed. New initial models that can be retrained (e.g. INCEPTION v.3 and v.4) have also been

developed (we used INCEPTION v.2). Although we explored some of these algorithms and initial models (see Methods S1), we did not optimize them because of the significant computational complexity in just optimizing Faster R-CNN/INCEPTION v.2 for fitness traits. Thus, in future studies, these algorithms and initial models should be more thoroughly explored to further improve fitness trait phenotyping.

We should emphasize that pictures of seeds or fruits are taken for record keeping and documentation purposes regardless of whether a machine vision-based approach or manual counting is used. After the picture is available, it takes our Faster R-CNNbased models c. 109 and 40 s to provide counts for a seed and fruit picture, respectively. By contrast, manual counting takes us c. 50 s per 100 seeds and 40 s per 100 fruits. Thus, as the seed and fruit number increases, our Faster R-CNN-based models have an even bigger advantage over manual counting.

By examining fitness traits, especially seed counts, we were able to observe phenotypic changes in loss-of-function mutants that were previously not detectable (pap2, Sun et al., 2012) or not reported (kin7.4 and hon5). In our relatively small sample of 12 mutants, effects on fitness were observed for three (25%). A similar percentage of lines with lower fitness than WT was reported by Rutter et al. (2017), who investigated the fitness effects of Arabidopsis T-DNA insertion lines using fruit number as a measure. They also found that a sizable percentage of lines had increased fitness compared with WT (12%), leading them to conclude that genetic redundancy is not common. We found that fruit counts could reveal fitness effects for two of three genes, indicating that seed counts are a better measure of fitness in some cases, such as when a genotype produces more fruits with fewer seeds per fruit. We are currently measuring both seed and fruit counts for a large number (>400) of mutants, which will allow us to obtain a more complete picture of the relative importance of fruit and seed counts for assessing fitness.

The seed counting pipeline that we established does not measure seed size, which is an agriculturally important trait associated with yield and seed viability (Sundaresan, 2005). By measuring seed weights, we found that *pap2* produces larger seeds than WT. Although we observed no clear difference in viability between them, seed size is a useful distinguishing characteristic between these genotypes. It might also provide insight into the underlying biology. For example, one possible reason for the increased seed size in the mutant is a lower fertilization rate, which would lead to fewer seeds and less restriction on seed growth (Herridge *et al.*, 2011; Fatihi *et al.*, 2013). Because measuring seed weights is time-consuming, a focus of our future work will be to adapt our pipeline to include approaches to measure seed size and number simultaneously.

Taken together, our results illustrate the importance of fitness traits in the study of gene functions and show that Faster R-CNN-based models, which can almost perfectly detect and count Arabidopsis seeds and also detect fruits with high accuracy, are valuable tools in large-scale studies of plant fitness. In future, we will use these tools to measure the fitness traits of a larger number of mutants to obtain a more complete picture of the effects of loss of gene function on fitness.



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Author contributions

PW, FM, JKC, PJK, MDL-S and S-HS conceived and designed the study. PW, FM, PD, SH, NLP, EV, EW, JKC, PJK and MDL-S performed data collection and analysis. PW, FM, MDL-S and S-HS wrote the manuscript. All authors read and approved the final manuscript. PW and FM are joint first authors.

ORCID

Jeffrey K. Conner (D https://orcid.org/0000-0003-1613-5826 Patrick J. Krysan (D https://orcid.org/0000-0003-4916-915X Melissa D. Lehti-Shiu (D https://orcid.org/0000-0003-1985-2687

Fanrui Meng b https://orcid.org/0000-0002-7911-6991 Nicholas L. Panchy b https://orcid.org/0000-0002-1551-3517 Shin-Han Shiu b https://orcid.org/0000-0001-6470-235X Peipei Wang b https://orcid.org/0000-0002-7580-9627 Eamon Winship b https://orcid.org/0000-0002-4200-3694

Data availability

All the scripts used in this study and the final seed and fruit counting models are available on GitHub at: https://github.com/ShiuLab/Manuscript_Code/tree/master/2022_Arabidopsis_seed_ and_fruit_count.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 The architecture of Faster Region-based CNN (R-CNN).

Fig. S2 Hyperparameter tuning for seed counting models.

Fig. S3 Computational efficiency of seed counting models.

Fig. S4 Example false negatives from the segmentation method using IMAGEJ and Faster Region-based CNN (R-CNN) models.

Fig. S5 Example images with different seed density index (SDI) values.

Fig. S6 Effect of seed density on the performance of the Faster Region-based CNN (R-CNN) models using different measures of performance.

Fig. S7 Hyperparameter tuning for fruit counting models.

Fig. S8 Fitness measurements for T-DNA insertion mutants of 12 genes.

Fig. S9 The proportion of fruits produced by 12 mutants that are shattered or green.

Methods S1 Plant growth conditions, seed image processing and seed counting.

Table S1 Lines used for training seed counting models.

Table S2 Lines used for training fruit counting models.

Table S3 Lines used for analysis of fitness.

Table S4 Fitness data for pap2, pap9, hon4, hon5, eif4b1 andeif4b2.

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