A flow cytometric method for estimating S-phase duration in plants

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Abstract

The duration of the DNA synthesis stage (S phase) of the cell cycle is fundamental in our understanding of cell cycle kinetics, cell proliferation, and DNA replication timing programs. Most S-phase duration estimates that exist for plants are based on indirect measurements. We present a method for directly estimating S-phase duration by pulse-labeling root tips or actively dividing suspension cells with the halogenated thymidine analog 5-ethynl-2′-deoxyuridine (EdU) and analyzing the time course of replication with bivariate flow cytometry. The transition between G₁ and G₂ DNA contents can be followed by measuring the mean DNA content of EdU-labeled S-phase nuclei as a function of time after the labeling pulse. We applied this technique to intact root tips of maize (Zea mays L.), rice (Oryza sativa L.), barley (Hordeum vulgare L.), and wheat (Triticum aestivum L.), and to actively dividing cell cultures of Arabidopsis (Arabidopsis thaliana (L.) Heynh.) and rice. Estimates of S-phase duration in root tips were remarkably consistent, varying only by ~3-fold, although the genome sizes of the species analyzed varied >40-fold.

Key words: Arabidopsis, barley, EdU, flow cytometry, maize, replication timing, rice, S-phase duration, wheat.

Introduction

Historically, most estimates of the duration of S phase and the cell cycle in plants have been made by analyzing the percentage of labeled mitoses (PLM) as a function of time after pulse-labeling with a DNA precursor. This technique was first described in mouse epithelium (Quastler and Sherman, 1959) and was the standard method of cell cycle analysis for nearly three decades (Van’t Hof, 1974; Grief et al., 2002; Francis et al., 2008, and references therein). Other methods that have been used to estimate S-phase duration in plants include double labeling (Wimber and Quastler, 1963), cell synchronization (Nagata et al., 1992; Lee et al., 1996; Cools et al., 2010), measurement of cell doubling time (Richard et al., 2001; Menges et al., 2006), kinetic analysis (Dhondt et al., 2010), and analysis of 5-ethynl-2′-deoxyuridine (EdU) labeling kinetics (Hayashi et al., 2013; Hu et al., 2015).

PLM (reviewed in Webster and Macleod, 1980; Shackney and Ritch, 1987), double labeling, and EdU labeling kinetics are limited logistically to relatively small numbers of cells. When implemented with radioactive DNA precursors (¹H and/or ¹⁴C), as they were originally, PLM and double labeling...
methods required lengthy autoradiographic exposure times and could have direct radiological effects on the cell cycle (De La Torre and Clowes, 1974; Gould and King, 1984; Darzynkiewicz et al., 1987; Fiorani and Beemster, 2006). Synchronization procedures necessarily perturb the normal cell cycle (Lee et al., 1996; Dolezel et al., 1999; Planchais et al., 2000; Menges and Murray, 2002; Cools et al., 2010). Moreover, the degree of synchronization depends on the individual characteristics of a cell population (Planchais et al., 2000) and is rarely complete (Cooper, 2003), particularly in roots (Lee et al., 1996). Methods based on estimates of cell doubling time are applicable mainly to isolated cells in suspension culture where, ideally, cells types are uniform and cell number and cell volume are proportional. However, a complication is the necessary assumption that all cells in suspension are cycling (Gould and King, 1984) when, in many cases, a substantial subpopulation is no longer dividing. In intact meristems, this approach is particularly complicated because meristems consist of multiple tissues and the proportionality of cell division and cell volume is generally not observed (Green, 1976; Fiorani and Beemster, 2006). Kinematic analysis requires a lengthy series of microscopic observations and complex mathematical analysis (Fiorani and Beemster, 2006; Rymen et al., 2010).

These methods have yielded many novel and informative analyses of the cell cycle but, in many cases, S-phase duration was inferred from measurements of other parameters. Flow cytometry is an excellent method for detailed analysis of proliferating cell populations as it provides the distribution of nuclei based on their DNA content (Fiorani and Beemster, 2006; Darzynkiewicz et al., 2011; Darzynkiewicz and Zhao, 2014). Additionally, it allows for rapid measurements of large cell populations, and the application of flow cytometry to plant systems was greatly facilitated by the development of a method for isolating and analyzing suspensions of isolated nuclei (Galbraith et al., 1983). Furthermore, bivariate flow cytometric analysis has the advantage of directly measuring the increase in DNA content of a cohort of labeled (replicating) cells as they move through S phase. Bivariate flow cytometry using the halogenated DNA precursor, 5-bromo-2’-deoxyuridine (BrdU), to label cells in S phase has been successfully applied to both animal (Reviewed in Darzynkiewicz et al., 2011) and plant systems (Lucretti et al., 1999) and was used to elucidate the replication timing program in Arabidopsis (Lee et al., 2010). When EdU is used as the DNA precursor in place of BrdU, visualization no longer requires denaturation of the DNA or cumbersome immunostaining procedures (Kotogany et al., 2010). EdU labeling, which can be detected by the covalent attachment of Alexa Fluor 488 (AF 488) via Click chemistry (Rostovtsev et al., 2002), has been used successfully for flow sorting and microscopic analysis of nuclei obtained from both suspension cultures and intact root tips (Bass et al., 2014, 2015; Wear et al., 2016).

To measure S-phase duration directly, we used a pulse-chase protocol to label DNA with EdU in a cohort of replicating cells. After various chase periods, we extracted nuclei and used bivariate flow cytometry to follow increases in the DNA content of the labeled cohort as a function of time.

We estimated the average S-phase durations in the root meristems of maize (Zea mays L.), rice (Oryza sativa L.), barley (Hordeum vulgare L.), and wheat (Triticum aestivum L.), and in Arabidopsis (Arabidopsis thaliana (L.) Heynh.) and cell悬 cell suspensions cultures. We found that S-phase duration in grass root tips is remarkably consistent, varying by just over 3-fold in species whose genome sizes span a nearly 40-fold range. When we compare the S-phase durations of rice root tip cells with those of cultured rice cells, we found that the cultured cells take approximately twice as long to complete DNA replication.

**Materials and methods**

**Plant growth**

Seed of Z. mays L. cultivar B73 provided by Mark Milliard (GRIN NPGS, North Central Regional Plant Introduction Station, Department of Agronomy, Iowa State University, Ames, IA, USA) were increased through one generation by 27 Farms of Homestead, Inc. (Homestead, FL, USA). Seed of O. sativa L. cultivar Nipponbare were provided by Dr Rongda Qu (Department of Crop Science, North Carolina State University, Raleigh, NC, USA). Seed of H. vulgare L. cultivar Morex were provided by Dr Kevin Smith (Department of Agronomy and Plant Genetics, University of Minnesota, St. Paul, MN, USA). Seed of T. aestivum L. cultivar Chinese Spring were provided by Dr Gina Brown-Guidera (Department of Crop Science, USDA, North Carolina State University, Raleigh, NC, USA) and Jon Raupp (Wheat Genetics Resource Center, Department of Plant Pathology, Kansas State University, Manhattan, KS, USA). Wild-type seed of A. thaliana (L.) Heynh. Col-0 were provided by Mary Dallas (Department of Plant and Microbial Biology, North Carolina State University, Raleigh, NC, USA).

For grass species, 50–200 seeds were used for each time point per species. The number of seeds remained consistent among time points and biological replicates for a given species. Maize seeds were imbibed overnight in sterile, distilled water with stirring and aeration prior to surface sterilization. Maize and de-hulled rice seeds were surface sterilized in a 10% commercial bleach solution containing 0.05% Tween-20 for 15 min with rotary mixing and washed 3–4 times with 2 vols of sterile water prior to germination. Twelve seeds were placed in sterile magenta boxes equipped with paper towels pre-wetted with 10 mL of sterile water. Seeds were germinated under constant, fluorescent dim light (6.75 μmol photons m⁻² s⁻¹) until primary roots were 2.5–4 cm long, which took 3 d for maize at 28 °C, 2 d for barley at 28 °C, 4 d for rice at 28 °C, and 3 d for wheat at 23 °C.

For Arabidopsis, 6000 seeds were used per time point. Arabidopsis seeds were surface sterilized in absolute ethanol for 5 min, followed by 20% commercial bleach containing 0.05% Tween-20 for 15–20 min with end-over-end mixing and washed five times with 1 vol. of sterile distilled water. Seeds were stored in sterile water and vernalized in the dark at 4 °C for 72 h. Three thousand seeds were germinated in rows per sterilized hydroponic dish (Alatorre-Cobos et al., 2014) equipped with 300 μm mesh in 250 mL of liquid Murashige and Skoog medium under constant fluorescent light (25.65 μmol photons m⁻² s⁻¹) until roots were 3–5 mm long, which took 4 d at 23 °C.

**Cell cultures**

An Arabidopsis (Col-0) cell line (Tanurdzic et al., 2008; Lee et al., 2010) was maintained in Gamborg’s B5 basal medium with minor salts (Sigma G5893) supplemented with 1.1 mg l⁻¹ 2,4-dichlorophenoxy acetic acid (2,4-D), 3 mM MES, and 3% sucrose. Cells were grown on a rotary shaker at 160 rpm under constant fluorescent light (24.3 μmol photons m⁻² s⁻¹) at 23 °C and subcultured every 7 d using a 1:8 dilution of inoculum into fresh medium.
A rice (O. sativa L.) cell line (Japonica cultivar Nipponbare; Lee et al., 2004) was maintained in AA medium (Thompson et al., 1986) supplemented with 2 mg l⁻¹ 2,4-D, 3 mM MES, and 3% sucrose. Cells were grown on a rotary shaker at 160 rpm in the dark at 27 °C and subcultured every 7 d using a 1:5 dilution of inoculum into fresh medium containing 0.005% pectinase.

**EdU labeling and fixation**

Roots were pulse-labeled as previously described by Wear et al. (2016). Roots of intact seedlings were rinsed in sterile distilled water and then incubated for 30 min in sterile water containing 25 μM EdU at 23 °C (wheat) or 28 °C (maize, rice, and barley) or 10 μM EdU at 23 °C (Arabidopsis roots). Incubations were carried out on a rotary shaker set to 65 rpm. The roots were rinsed twice with 2–3 vols of sterile water and the EdU label was chased for various times with 25 μM (maize and rice), 100 μM (maize, rice, barley, wheat, and Arabidopsis roots), or 200 μM (wheat) thymidine prepared in sterile water. Chase conditions using varying thymidine concentrations were used to resolve the appearance of the residually labeled ‘arm’ of nuclei observed in the flow cytograms of maize, rice, and wheat (Fig. 1A; Supplementary Figs S1, S2A, C at JXB online and described in detail in the Results). Roots were then rinsed twice with 2–3 vols of sterile water. For grass species, terminal 1 mm (maize) or 5 mm (rice, barley, and wheat) root segments were excised and fixed in 1% formaldehyde in 1× phosphate-buffered saline (PBS) or 5 min under vacuum. The formaldehyde reaction was quenched by adding glycine to a final concentration of 0.125 M and incubating at room temperature with end-over-end mixing for 5 min. The formaldehyde reaction was quenched by adding glycine to a final concentration of 0.125 M for 5 min under vacuum. Fixed root tips were washed three times with 1 vol. of 1× PBS, snap-frozen in liquid nitrogen, and stored at −70 °C until preparation of nuclei. For Arabidopsis roots, intact roots were fixed, quenched, and washed with the same conditions as described above and then terminal root segments ranging from 3 mm to 5 mm long were excised and snap-frozen as above.

Arabidopsis and rice cell suspensions were pulse-labeled as described by Wear et al. (2016). Four (Arabidopsis) or three (rice) days after transfer, logarithmically growing cells from a 50 ml culture were labeled by adding 12.5 μl or 31.25 μl of a 40 mM solution of EdU prepared in DMSO to the culture medium to a final concentration of 10 μM (Arabidopsis) or 25 μM EdU (rice), respectively. An EdU pulse concentration of 10 μM for Arabidopsis cells was used to maintain consistency with concentrations used in replication timing studies, and an EdU pulse concentration of 25 μM for rice cells was used to maintain consistency with concentrations used in labeling intact rice root meristems. Cells were incubated for 30 min at 23 °C (Arabidopsis) or 27 °C (rice) on a rotary shaker set to 160 rpm. The EdU was chased for various times by adding 125 μl of a 40 mM solution of thymidine in sterile water to the labeled cultures to a final concentration of 100 μM. For each time point after the chase, 10 ml of culture was removed and centrifuged at 400 g for 2 min. The supernatant was removed and cells were fixed by re-suspending the pellet in 1% formaldehyde in 1× PBS and incubating at room temperature with end-over-end mixing for 15 min. The formaldehyde reaction was quenched by adding glycine to a final concentration of 0.125 M and incubating at room temperature with end-over-end mixing for 5 min. The fixed cells were pelleted by centrifugation at 400 g for 2 min and the fixative/quench solution was discarded. The cell pellet was re-suspended and washed three times in 1 vol. of 1× PBS with centrifugation as above between each wash step. After the final wash step, cells were pelleted by centrifugation as above and the PBS was discarded. The cell pellet was snap-frozen in liquid nitrogen and stored at −70 °C until preparation of nuclei.

**Fig. 1.** Flow cytometric analysis of an S-phase duration time course in maize root tip cells. Roots of 3-day-old maize seedlings were pulse-labeled with 25 μM EdU for 30 min, washed, and then chased with 25 μM thymidine for select hourly intervals. Terminal 1 mm segments of the root tips were excised, fixed, and frozen. Nuclei were prepared and analyzed by flow cytometry. (A) Bivariate flow cytograms of EdU incorporation (AF 488 fluorescence) and relative DNA content (DAPI fluorescence). The flow cytograms show labeled S-phase populations (F₁) and unlabeled G₁ (F₂) and G₂/M (F₃) populations at various times during the chase. The mean DNA content for each population was determined by calculating the mean DAPI fluorescence within the gated region (red boxes for EdU-labeled S-phase nuclei and black boxes for unlabeled G₁ and G₂/M nuclei). At 0 h, the gate used to calculate mean DAPI fluorescence for labeled nuclei included the entire, main S-phase arc (red box). At time points after 0 h, the gate used to calculate mean DAPI fluorescence of labeled nuclei (smaller red boxes in 2–7 h) excluded the labeled nuclei that had divided and returned to G₁ (gray boxes directly above unlabeled G₁ and juxtaposed to red boxes in flow cytograms). (B) Corresponding univariate histograms of relative DNA content (DAPI fluorescence) of EdU-labeled, S-phase nuclei. The S-phase histograms show the distribution of the main S-phase nuclei (including those that had returned to G₁) immediately after labeling and throughout the chase and have been standardized so that each time point contains the same number of total events. The bivariate flow cytograms and corresponding histograms represent a single biological replicate.
Isolation of nuclei

Nuclei were isolated by a modification of the chopping method (Galbraith et al., 1983) as described by Wear et al. (2016). Fixed and frozen root tips or cell pellets were ground at 4 °C in 25–35 ml of cell lysis buffer (CLB: 15 mM Tris, pH 7.5, 2 mM Na3EDTA, 80 mM KC1, 20 mM NaCl, 0.1% Triton X-100, and 15 mM 2-mercaptoethanol) using the lowest setting on a commercial food processor (Cuisinart® Mini-Prep® Processor, model DLC-1SS). The ground suspension was incubated at 4 °C for 5 min with gentle swirling half way through the incubation to maximize the yield of nuclei and then filtered through a double layer of Miracloth. The filtrate was centrifuged at 400 g for 5 min and the pellet of nuclei was re-suspended in 1 ml of modified CLB without EDTA and 2-mercaptoethanol.

Click chemistry and DAPI staining

Nuclei were centrifuged at 200 g, 4 °C, for 5 min and the supernatant was discarded. Labeled nuclei were thoroughly re-suspended and coupled to AF 488 in 0.5 ml of Click-iT® reaction buffer according to the manufacturer’s instructions (Click-iT® EdU Alexa Fluor® 488 Imaging Kit, Life Technologies). Nuclei were then centrifuged as before, washed with 2 vols of CLB, and re-suspended in 0.5–1.0 ml of CLB containing 2 μg ml⁻¹ DAPI. Following each centrifugation step, the nuclei were carefully and thoroughly re-suspended to minimize aggregation. The final suspension was filtered through a CellTrics 20 μm nylon filter (Partec) prior to flow cytometric analysis.

Flow cytometric analysis

Nuclei were analyzed with an InFlux (BD Biosciences) cell analyzer/sorter equipped with a 355 nm UV laser, a 488 nm sapphire laser, and BD FACSTM™ Software v. 1.0.0.650, using 1× PBS as the sheath buffer. The software recorded forward scatter (FSC), side scatter (SSC), emission at 460/50 nm (DAPI fluorescence), and emission at 530/40 nm (AF 488 fluorescence). FSC was used as the trigger and to set the event threshold. Dot plots of FSC versus SSC, FSC versus DAPI fluorescence, and SSC versus DAPI fluorescence were used to locate and gate populations based on light scattering properties and DNA content (described in detail in Wear et al., 2016). A bivariate dot plot of AF 488 fluorescence (log scale) versus DAPI fluorescence (linear scale) was created to visualize labeled and unlabeled nuclei at each time point in the experiment. In the plots, labeled nuclei at different stages of S phase formed an arc above the unlabeled G₀ and G₁ populations of nuclei.

Relative movement

FlowJo v. 10.0.6 (Tree Star, Inc.) was used for flow cytometric data analysis. For each experiment, we analyzed at least three biological replicates conducted at different times with different batches of plant material. Within each biological replicate, we analyzed a minimum of three separate data files per time point. SSC versus DAPI plots were used to determine appropriate gates to exclude small debris from the analysis (described in detail in Wear et al., 2016). AF 488 versus DAPI plots were used to delineate gates for labeled nuclei in S phase and unlabeled nuclei in G₀ or G₂/M, using similar gates for each of the biological replicates for a given species. The mean DAPI fluorescence was quantified for each of the S-phase, G₀, and G₂/M gated populations. At 0 h (i.e. immediately after the EdU pulse-label), the gate included to calculate mean DAPI fluorescence for labeled, undivided nuclei included the entirety of the main S-phase arc (Fig. 1A, 0 h, red gate). At all time points after 0 h, the gate used to calculate mean DAPI fluorescence of labeled nuclei was drawn such that it excluded nuclei that had divided and returned to G₀ (Fig. 1A, 2–7 h, smaller red gates). For each time point, the relative movement (RM) of EdU-labeled nuclei in S phase was calculated by comparing their mean DAPI fluorescence relative to the mean DAPI fluorescence of the G₀ and G₂/M populations using the equation: RM=(F₀–F₂G/M)/(F₀–F₂G₀) (Begg et al., 1985), where F₀=mean DAPI fluorescence of EdU-labeled nuclei, F₂G₀=mean DAPI fluorescence of unlabeled G₀ nuclei, and F₂G/M=mean DAPI fluorescence of unlabeled G₂/M nuclei. For each species, the average RM values from multiple biological replicates were combined and plotted as a function of time.

S-phase duration

S-phase duration estimates were calculated from statistical analyses of an RM plot of combined biological replicates using R v. 3.2.3 loaded with the package ‘Segmented’ v. 0.5–1.4 (Muggeo, 2008). A generalized linear model was first applied to the data points, and the Davies test was used to test for a non-constant slope parameter. ‘Segmented’ was used to estimate a regression model with a piecewise linear relationship having a single breakpoint and two lines. The 95% confidence intervals were computed for the slope of each of the two fitted regression lines and an R² value was calculated. Lower and upper estimates of S-phase duration were derived from the estimated breakpoint and extrapolation of the first component line (line below the estimated breakpoint) to a theoretical RM value of 1.0.

Results

Flow cytometric analysis of EdU-labeled S-phase nuclei

Seedlings or cell suspension cultures were pulse-labeled with EdU and then chased with excess thymidine for various periods of time as described in the Materials and methods. At each time point, root segments or cells were fixed, nuclei were isolated, the incorporated EdU was coupled to AF 488, and total DNA was stained with DAPI. We analyzed the nuclei by flow cytometry to produce bivariate plots of relative DNA content (DAPI fluorescence) and EdU incorporation (AF 488 fluorescence) as well as univariate plots showing the distribution of relative DNA contents in the cohort of EdU-labeled (replicating) nuclei. Examples of such plots for nuclei from maize root tips are shown in Fig. 1A, B.

Three distinct populations of nuclei were identified in the bivariate plots—unlabeled nuclei in G₀, with 2C DNA content, unlabeled nuclei in G₂/M with 4C DNA content, and EdU-labeled S-phase nuclei with DNA contents between 2C and 4C. At the end of the labeling period (Fig. 1A, 0 h), the DNA content of S-phase nuclei was intermediate between the two extremes and formed a distinct arc above the unlabeled populations in the flow cytograms. Samples harvested at various times during the thymidine chase were used to track the movement of EdU-labeled nuclei as they progressed through S phase by following increases in DNA content of the nuclei in the red box (Fig. 1A). A subset of labeled S-phase nuclei that were probably near the end of S phase when the pulse label began, divided and re-entered G₀, creating a new group of labeled nuclei with 2C DNA content (Fig. 1A; 4–7 h, gray boxes directly above unlabeled G₀). The distribution of the main EdU-labeled nuclei population was visualized in histograms for each time point (Fig. 1B, red box at 0 h and gray box plus red box at 2–7 h) and confirmed the even distribution of DNA content between 2C and 4C at the end of the labeling period (Fig. 1B, 0 h), the movement of the main cohort of labeled nuclei (Fig. 1B, after 0 h), and the return of labeled nuclei with 2C DNA content (Fig. 1B, 4–7 h).
An ‘arm’ of less intensively labeled nuclei (Fig. 1A, 2–4h) was observed below the S-phase gate and extended diagonally upwards from 2C across to 4C. These nuclei progressed towards 4C but remained below the fully labeled gated 4C nuclei, eventually forming a downward extension of the S-phase arc (Fig. 1A, 5–7h). Similar ‘arms’ were observed in the flow cytograms of rice and wheat roots (see Supplementary Fig. S2A, C at JXB online) and Arabidopsis cell culture (Fig. 4A). The arms most probably reflect a low level of EdU labeling that continued after the start of the chase period. In cell cultures, the chase was initiated by adding excess thymidine, without washing away the EdU. In the case of root tips, after the pulse-label was complete, excess EdU was washed away from the root tips before placing them in the thymidine solution. However, it is likely that some EdU remained in intercellular spaces and intracellular pools. In both cases, it may have taken some time for the thymidine chase to become fully effective, and the effectiveness of the chase may have differed between species or cell types. For example, increasing the concentration of the thymidine chase from 25 μM to 100 μM significantly decreased the number of nuclei in the ‘arm’ for maize root tip samples (Supplementary Fig. S1), but residual incorporation occurred even with a chase concentration as high as 200 μM in wheat root tips (Supplementary Fig. S2C). It is important to note that EdU labeling was displayed on a logarithmic scale in the flow cytograms and most of the residual labeling was 10- to 100-fold lower than that of fully labeled nuclei. Furthermore, the residually labeled nuclei were excluded from the gates used to follow the DNA content of pulse-labeled nuclei, and did not affect the RM measurements described below or the resulting estimates of S-phase duration.

Relative movement and measurement of S-phase duration

To quantify progression through S phase, we drew gates around the three populations of G1, G/M, and S-phase (EdU-labeled) nuclei and calculated the mean DAPI fluorescence, a measure of DNA content, for each population. The RM of the nuclei in S phase was then calculated for each time point (refer to the Materials and methods for gating and RM calculations). Assuming that the increase in DNA content occurred at a constant rate throughout S phase, the average DNA content of labeled nuclei at 0 h should be half way between 2C and 4C, with an RM of 0.5 (Begg et al., 1985). In theory, the RM should continue to increase as nuclei progress through S phase, with a value of 1.0 being reached when all nuclei that initiated DNA synthesis during the labeling period finished replication (Begg et al., 1985).

In our experiments, RM values at 0 h (samples taken immediately after the 30 min labeling period) ranged from 0.45 to 0.61, essentially as predicted (Figs 2, 3D–F; 4E; F; 0h). RM values also increased over time as expected, although, in the actively growing systems we studied, they did not reach the theoretical value of 1.0. This value would only have been reached if the labeled cells all accumulated in G2 without returning to G1 during the time course of the experiment, as apparently was the case in the experiments of Begg and colleagues with BrdU-labeled CHO cells (Begg et al., 1985). In our calculations, we minimized the effect of G1 return by excluding EdU-labeled nuclei with 2C DNA content (Fig. 1A; 2–7 h, gray boxes). However, labeled nuclei that failed to complete S phase or that returned to G1 and subsequently began a second round of DNA synthesis caused the RM values to plateau at an RM value <1.0.

To estimate the point at which a plateau was reached, we applied a segmented linear regression analysis in which two linear components and a single breakpoint were fit to the data. To ensure high confidence fitting of the linear regression, data from multiple biological replicates were combined prior to statistical analysis. The precision of the analysis was indicated by 95% confidence intervals on the slopes of each of the component lines (Figs 2, 3D–F; 4E, F). Two estimates of S-phase duration were obtained from this analysis. One estimate involved taking the slope of the first linear component as the initial rate of DNA replication and extrapolating the line to an RM value of 1.0. A second estimation
procedure assumed that the breakpoint in the RM curve represents essentially complete replication of the predominant cell population. The breakpoint estimate probably best reflects the behavior of the majority of cells with labeled nuclei because it minimizes the effect of cells that failed to progress through S phase, as well as the effect of cells that returned to G1 and began a second round of synthesis. We cannot formally distinguish between these two possibilities, which are not mutually exclusive. However, a second round of DNA synthesis in a subset of the labeled cells seems quite likely in a rapidly dividing population such as the root tips used here. Initiating a second round of DNA synthesis within the time frame of our experiments would require G1 to be traversed quite rapidly, but brevity of G1, or even its absence, is well documented in rapidly dividing animal cells (White and Dalton, 2005; Orford and Scadden, 2008), and G1 durations between 0.5h and 1.0h have been documented in root meristems of monocots (Evans and Rees, 1971).

The following paragraphs describe our analyses for several different species. S-phase duration estimates and RM values at the breakpoint for each species studied are summarized in Table 1.

S-phase duration in root tips

Immediately after EdU labeling, S-phase nuclei from maize root tips were distributed evenly between 2C and 4C (Fig. 1A, B; 0h). Through 4h, there was a steady increase in the amount of labeled nuclei that reached 4C. At some point between 3h and 4h, labeled nuclei began to return to 2C. At subsequent time points, the number of labeled nuclei at 2C steadily increased and those at 4C slowly decreased. As discussed above, it is likely that some of these returned, labeled nuclei initiated a second round of DNA synthesis, resulting in a steady-state population of labeled nuclei with intermediate DNA content and lower the RM value at the breakpoint (0.85) (Fig. 2; Table 1). The effect was to increase the difference between the breakpoint (2.7h) and extrapolated (3.9h) estimates of S-phase duration (Fig. 2). These observations suggested that a large fraction of maize root tip nuclei completed S phase in ~2.7h.

Histograms from time course analyses with root tips of other grass species are presented in Fig. 3 (see Supplementary Fig. S2 for flow cytograms). In rice (Fig. 3A), a large fraction of labeled nuclei approached 4C after a 1h chase period. Between 1h and 2h, a small fraction of labeled nuclei returned to 2C and the fraction of labeled nuclei in 2C significantly increased between 2h and 3h. Like maize, the rice RM value at the breakpoint, 0.84 (Fig 3D; Table 1), was relatively low, most probably because of labeled nuclei undergoing a second round of replication. As a consequence, the two estimates of S-phase duration (1.2h and 1.9h) were different. We suggest that the largest cohort of rice root tip nuclei completed replication in 1.2h (Fig. 3D).

In barley (Fig. 3B), a large fraction of the labeled nuclei reached 4C by 3h and return of labeled nuclei to 2C was apparent between 2h and 3h. A significant fraction of labeled nuclei had returned to 2C by 4h. Some of these may have undergone a second round of replication, but the effect was smaller than in maize or rice. Thus, the RM value at the breakpoint, 0.94 (Fig. 3E; Table 1), was closer to 1.0 and the breakpoint and extrapolation estimates of S-phase duration, 2.3h and 2.7h, respectively, were more similar (Fig. 3E).

In wheat (Fig. 3C), a large population of labeled nuclei reached 4C at ~2–3h and a substantial fraction returned to 2C by 3h. As in barley, the RM value at the breakpoint, 0.94 (Fig. 3F; Table 1), was high and the two estimates of S-phase duration, 2.5h and 2.9h, were similar (Fig. 3F).

S-phase duration in suspension cultures

In Arabidopsis cultured cells (Fig. 4A, B), a large fraction of labeled nuclei reached 4C between 1h and 2h. However, the return of labeled nuclei to 2C occurred between 4h and 6h, in contrast to 2–4h in the root tip samples, and probably accounted for the decreased numbers of labeled nuclei with intermediate DNA contents in the flow cytograms (Fig. 4A). This observation is not too surprising given that cultured plant cells often have an extended cell cycle duration (refer to table 1 in Gould and King, 1984; and table 8 in Griffiths et al., 2002) with G1 duration considerably longer than that of root tips (Gould and King, 1984). Returning nuclei, therefore, had a smaller influence on RM calculations, although there may have been a small number of nuclei progressing slowly or perhaps even arrested in mid-S phase. The RM value for the Arabidopsis cells at the breakpoint, 0.91 (Fig. 4E; Table 1), was high and both 1.5h and 1.9h are good estimates of S phase for the largest cohort of proliferating nuclei (Fig. 4E).

In rice cell culture, a large cohort of labeled nuclei reached 4C by 3h (Fig. 4C, D) and a significant return to 2C occurred

| Species                  | Tissue type | S-phase duration breakpoint/extrapolated (h) | RM value at breakpoint | Genome size (Gbp) |
|--------------------------|-------------|---------------------------------------------|------------------------|-------------------|
| Maize cv. B73            | Roots       | 2.7/3.9                                     | 0.85                   | 2.3 (Schnable et al., 2009) |
| Barley cv. Morex         | Roots       | 2.3/2.7                                     | 0.94                   | 5.1 (Mayer et al., 2012) |
| Wheat cv. Chinese Spring | Roots       | 2.5/2.9                                     | 0.94                   | 17 (Brenchley et al., 2012) |
| Rice cv. Nipponbare      | Roots       | 1.2/1.9                                     | 0.84                   | 0.39 (Matsumoto et al., 2005) |
| Rice cv. Nipponbare      | Culture     | 2.6/4.7                                     | 0.81                   | 0.39 (Matsumoto et al., 2005)* |
| Arabidopsis (Col 0)      | Culture     | 1.5/1.9                                     | 0.91                   | 0.13 (Arabidopsis Genome Initiative, 2000)* |

* Genome sizes are known to vary in tissue culture (Lee and Phillips, 1988).
between 3 h and 4 h. Notably, a large subpopulation of nuclei either appeared to arrest part way through S phase or proceeded much more slowly than the main cohort of labeled nuclei. This subpopulation is clearly visible in the flow cytograms at 3–5 h and is visualized as a pronounced shoulder on the 4C peak in the corresponding histograms (Fig. 4C, D). The subpopulation lowered the RM value at the breakpoint, 0.81 (Fig. 4F; Table 1), and resulted in a large difference between the two S-phase estimates, 2.6 h and 4.7 h (Fig. 4F). In this case, 4.7 h is probably a considerable overestimate of S-phase duration for the major class of labeled nuclei, and 2.6 h more accurately reflects the S-phase duration for this group.

Discussion

We sought a technique for directly estimating S-phase duration in large populations of cells without the need for disruptive synchronization treatments. Thus, we used bivariate flow cytometry to follow the increase in DNA mass that occurs between G1 and G2 using EdU incorporation as a tool to focus on a cohort of cells actively engaged in DNA synthesis. Our method can be applied to any system containing a sufficient number of cells that can be labeled with EdU and can be analyzed by flow cytometry as either cells or nuclei. In our hands, the method is highly reproducible and reduces the complexity of obtaining robust estimates of S-phase duration for large populations of cells.

The root tip system and EdU labeling techniques are applicable to a wide variety of plant species. Root tips efficiently take up both EdU and thymidine, and contain a large fraction of proliferating cells. Advantages of EdU labeling include the fact that visualization of incorporated EdU is achieved rapidly and under mild conditions that do not degrade nuclei or subnuclear structure (Kotogany et al., 2010). Our flow cytometric procedure
provides a direct measurement of the increase in DNA mass that occurs during replication and can easily be applied to unsynchronized cell populations. As such, it eliminates the necessity for disruptive synchronization treatments such as sucrose starvation or addition of chemical inhibitors (Dolezel et al., 1999; Planchais et al., 2000; Menges and Murray, 2002) and can be applied to nuclei derived from cell cultures as well as intact organs. Large numbers of nuclei can be analyzed rapidly, offering increased statistical power compared with techniques that require scoring of individual nuclei under the microscope. Microscopic techniques will continue to be useful in many instances, especially in cases where material is limited and/or where it is important to analyze S phase in an anatomically defined subset of cells. However, flow cytometry is advantageous in systems in which large numbers of cells or nuclei from proliferating cell populations can be isolated for analysis.

**Fig. 4.** Flow cytometric analysis and estimation of S-phase duration in Arabidopsis and rice suspension cell cultures. Four (Arabidopsis) or three (rice) days after transfer, cells in logarithmic growth phase were pulse-labeled with 10 μM (Arabidopsis) or 25 μM (rice) EdU for 30 min and then chased with 100 μM thymidine for select hourly intervals. Cells were pelleted, fixed, washed, and frozen. Nuclei were prepared and analyzed by flow cytometry as described in Fig. 1. (A–D) Bivariate flow cytograms and corresponding histograms of relative DNA content of S-phase nuclei isolated from cell suspension cultures of (A, B) Arabidopsis and (C, D) rice. Data represent single biological replicates. (E, F) RM plots for (E) Arabidopsis and (F) rice cell cultures. For each species, RM values from multiple biological replicates were calculated and plotted as a function of time. Segmented regression analysis was applied to the data of combined biological replicates as described in Fig. 2. The $R^2$ value of the segmented fit, slope of the first component line, and lower and upper average S-phase duration estimates are as follows: (E) Arabidopsis, $R^2=0.99$, $y=0.2326x+0.5654$ and 1.5 h/1.9 h; and (F) rice, $R^2=0.95$, $y=0.0926x+0.5676$ and 2.6 h/4.7 h.
Bivariate flow cytometric analysis of the cell cycle using BrdU has been used to follow DNA synthesis in both plant (Lucetti et al., 1999) and animal (Begg et al., 1985) systems. Begg and colleagues applied this technique to estimate the duration of S phase, using RM plots similar to ours. Under their conditions with mammalian cells, RM values approached 1.0 as the population of labeled cells completed a single round of replication. However, in our material, some labeled nuclei initiated a second round of DNA synthesis and, in some cases, a subset of nuclei progressed slowly or arrested prior to completion of S phase. The presence of one or both of these two types of intermediate DNA contents lowered RM values and prevented them from reaching 1.0 as was achieved in the mammalian systems. In all cases in this study, the RM plots showed a clear break in slope before reaching 1.0, and this led us to apply a segmented linear regression analysis to fit two components to our data and to determine the breakpoint between them. We interpreted the breakpoint as representing the time at which the largest cohort of nuclei completed DNA synthesis. In most cases, we think this is the best estimate of S-phase duration for the population under study, a view supported by detailed analysis of flow cytometry profiles presented in the Results.

**Cells versus roots**

Many features of the cell culture environment are known to affect total cell cycle length and it is well documented that suspension cell cultures typically have a much longer G1 phase than root meristems (Gould and King, 1984, and references therein). Less is known about the effect of culture parameters on the S-phase component of the cell cycle. The fact that S-phase length estimates vary as much as 2- to 3-fold between different studies using suspension cells from the same species suggests that S-phase progression is likely to be sensitive to multiple environmental influences (see references for Acer pseudoplatanus and Haplopappus gracilis in table 1 of Gould and King, 1984).

We compared S-phase duration for cultured cells and root meristems in both rice and Arabidopsis. Unfortunately, we could not obtain data of sufficient quality from Arabidopsis root tips (see Supplementary Fig. S3 at JXB online). However, comparison of rice suspension culture cells with rice root tip meristem cells showed that S phase is shorter in root tips. RM plots for suspension cells showed a lower slope of the first component line (compare Figs 4F and 3D), indicative of the presence of a slower replicating group of cells in addition to the main cohort. Extrapolating from the first component to estimate an overall average S phase for suspension cells and root tips gave values of 4.7h and 1.9h (Table 1), respectively, for a 2.5-fold difference. Estimates derived from breakpoint values, which reflect the most rapidly replicating major cohort of nuclei, differed by more than a factor of two (2.6h for suspension cells and 1.2h for root tips; Table 1).

The presence of a more slowly proliferating subset of cells in plant cell suspension cultures may reflect that many such cultures are comprised of clumps or multicellular aggregates alongside isolated, single cells. Cells in the interior of aggregates are subject to a different microenvironment compared with free floating cells or cells on the surface of aggregates (Street, 1973), including differences in oxygen (Hulst et al., 1989) and nutrient availability (Patil and Roberts, 2013). Other factors that may contribute to the difference in S-phase duration include genetic and epigenetic instability, such as chromosome rearrangements, DNA methylation, and somatic mutation (reviewed by Lee and Phillips, 1988; Phillips et al., 1994; Tanurdzic et al., 2008).

**Comparison with other reported estimates**

We compared our S-phase duration estimates with previously reported estimates for species included in our study (Supplementary Table S1). There is a clear tendency for our estimates to be shorter than those in previous reports. There are several explanations for the differences. Because flow cytograms do not cleanly resolve nuclei that have replicated 100% of their DNA from those having replicated slightly less than 100%, the breakpoint method may underestimate the full duration of S phase, especially in cases where a subset of the DNA (e.g. maize knob heterochromatin or B chromosomes) replicates extremely late in S phase. The extrapolation method is less sensitive to this problem but can significantly overestimate duration when a fraction of cells fail to complete S phase, as observed in our rice suspension culture. In other cases, the breakpoint and extrapolation methods may be thought of as yielding lower and upper estimates of S-phase duration, respectively.

Other reasons for variation in S-phase estimates include the fact that different subsets of cells within the root meristem can vary in S-phase duration (Barlow and Macdonald, 1973; Van’t Hof, 1974, and references therein; Hayashi et al., 2013). Some techniques focus on certain groups of cells, while our method provides an estimate of the average duration for the major cohort of cells in S phase. Temperature and a variety of other environmental variables are also known to affect S-phase duration (Van’t Hof, 1974, and references therein; Verma, 1980; Francis and Barlow, 1988; Grif et al., 2002). Older estimates may have been influenced by technical factors inherent to the methodology, such as radiological effects (De La Torre and Clowes, 1974; reviewed in Gould and King, 1984; Darzynkiewicz et al., 1987; Fiorani and Beemster, 2006) as well as sampling errors and/or mathematical assumptions (Shackney and Ritch, 1987; Grif et al., 2002). Considering the many factors that can affect the duration of S phase, it is evident that measurement of S should be made under conditions that closely match those of other experiments with which they will be compared. Our method was designed to match the experimental conditions we use to study replication timing programs but can accommodate a wide range of other conditions.

S-phase duration estimates in other eukaryotes exhibit similar within-species variability as has been observed for plants (Supplementary Table S2). More diverse cell types have been studied in mammalian systems such as mouse, rat, and human, whereas in plants most reports focus on roots. The broader sampling of tissue types and cell lines in mammals...
may explain the discovery of a few examples of notably long S-phase estimates in some specialized tissues or lines.

**Relationship to genome size**

We estimated the average S-phase duration in root tips from a selection of grass species with widely varying genome sizes to address the question as to whether S-phase duration was correlated to genome size (Table 1). The selected species cover a 40-fold range in genome size. However, comparison of the breakpoint estimates of S-phase duration only uncovered an ~2-fold difference in the duration of S phase. When both the breakpoint and extrapolation estimates of S-phase duration are considered, there is an ~3-fold difference in the duration of S phase. The lack of an obvious relationship of S-phase duration and genome size was initially surprising given early reports that used the PLM method to show a trend toward longer S phases in species with larger genome sizes (Van’t Hof, 1965; Evans and Rees, 1971). The significance of this trend depended heavily on the inclusion of species from such genera such as *Allium* and *Tradescantia* with very large genomes in the range of 20–33 Gbp. In contrast, plants with moderately sized genomes showed only small, apparently random, variations in S-phase duration. Similarly, the sampling of previously reported plant S-phase duration values in Supplementary Table S1 shows no correlation to genome size. Several studies (reviewed by Bennett, 1972; Francis et al., 2008) reported a strong dependence of total cell cycle time on genome size. However, given that DNA replication occurs in separate replions, many of which can replicate simultaneously, S-phase duration may not be strongly dependent on genome size.

**Supplementary data**

Supplementary data are available at *JXB* online

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**Figure S1.** Flow cytometric analysis of maize roots chased with 100 μM thymidine.

**Figure S2.** Flow cytometric analysis of rice, barley, and wheat root tips.

**Figure S3.** Hydroponic growth system, flow cytometric analysis, and DNA content histograms of S-phase nuclei from roots of 4-day-old Arabidopsis seedlings.

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**Table S1.** Comparison of S-phase duration estimates in plants.

**Table S2.** S-phase duration estimates in various eukaryotes.

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**References**

Alatorre-Cobos F, Calderon-Vazquez C, Ibarra-Laclette E, et al. 2014. An improved, low-cost, hydroponic system for growing Arabidopsis and other plant species under aseptic conditions. BMC Plant Biology 14, 69.

**Arabidopsis Genome Initiative.** 2000. Analysis of the genome sequence of the flowering plant Arabidopsis thaliana. Nature 408, 796–815.

**Barlow PW, Macdonald PDM.** 1973. Analysis of the mitotic cell cycle in the root meristem of *Zea mays*. Proceedings of the Royal Society B: Biological Sciences 183, 385–398.

**Bass HW, Hoffman GG, Lee TJ, Wear EE, Joseph SR, Allen GC, Hanley-Bowdoin L, Thompson WF.** 2015. Defining multiple, distinct, and shared spatiotemporal patterns of DNA replication and endoreduplication from 3D image analysis of developing maize (*Zea mays L.*) root tip nuclei. Plant Molecular Biology 89, 339–351.

**Bass HW, Wear EE, Lee TJ, Hoffman GG, Gumber HK, Allen GC, Thompson WF, Hanley-Bowdoin L.** 2014. A maize root tip system to study DNA replication programmes in somatic and endocycling nuclei during plant development. Journal of Experimental Botany 65, 2747–2756.

**Begg AC, McNally NJ, Shriève DC, Karcher H.** 1985. A method to measure the duration of DNA synthesis and the potential doubling time from a single sample. Cytometry 6, 620–626.

**Bennett MD.** 1972. Nuclear DNA content and minimum generation time in herbaceous plants. Proceedings of the Royal Society B: Biological Sciences 181, 109–115.

**Brenchley R, Spannagl M, Pfeifer M, et al.** 2012. Analysis of the bread wheat genome using whole-genome shotgun sequencing. Nature 491, 705–710.

**Cools T, Iantcheva A, Maes S, Van den Dale H, De Veylder L.** 2010. A replication stress-induced synchronization method for Arabidopsis thaliana root meristems. The Plant Journal 64, 705–714.

**Cooper S.** 2003. Rethinking synchronization of mammalian cells for cell cycle analysis. Cellular and Molecular Life Sciences 60, 1099–1106.

**Darzyynkiewicz Z, Traganos F, Kimmel M.** 1987. Assay of cell cycle kinetics by multivariate flow cytometry using the principle of stathmokinesis In: Gray JW, Darzyynkiewicz Z, eds. Techniques in cell cycle analysis. Humana Press: Clifton, NJ, 291–336.

**Darzyynkiewicz Z, Traganos F, Zhao H, Halicka HD, Li J.** 2011. Cytometry of DNA replication and RNA synthesis: historical perspective and recent advances based on ‘click chemistry’. Cytometry A 79, 328–337.

**Darzyynkiewicz Z, Zhao H.** 2014. Cell cycle analysis by flow cytometry. eLS.

**De La Torre C, Clowes FAL.** 1974. Thymidine and the measurement of rates of mitosis in meristems. New Phytologist 73, 919–925.

**Dhondt S, Coppons F, De Winter F, Swarup K, Merks RM, Inze D, Bennett MJ, Beemster GT.** 2010. SHORT-ROOT and SCAPECROW regulate leaf growth in Arabidopsis by stimulating S-phase progression of the cell cycle. Plant Physiology 154, 1183–1195.

**Dolezel J, Cihalíková J, Weiserova J, Lucretti S.** 1999. Cell cycle synchronization in plant root meristems. Methods in Cell Science 21, 95–107.

**Evans GM, Rees H.** 1971. Mitotic cycles in dicotyledons and monocotyledons. Nature 233, 350–351.

** Fiorani F, Beemster GT.** 2006. Quantitative analyses of cell division in plants. Plant Molecular Biology 60, 963–979.

**Francis D, Barlow PW.** 1988. Temperature and the cell cycle. Symposia of the Society for Experimental Biology 42, 181–201.

**Francis D, Davies MS, Barlow PW.** 2008. A strong nucleotide effect on the cell cycle regardless of ploidy level. Annals of Botany 101, 747–757.

**Galbraith DW, Harksen KR, Maddox JM, Ayres NM, Sharma DP, Firoozabady E.** 1983. Rapid flow cytometric analysis of the cell cycle in intact plant tissues. Science 220, 1049–1051.

**Gould AR, King PJ.** 1984. Control of the cell cycle in cultured plant cells. Critical Reviews in Plant Sciences 1, 315–344.

**Green PB.** 1976. Growth and cell pattern formation on an axis: critique of concepts, terminology and modes of study. Botanical Gazette 137, 187–202.

**Grif VG, Ivanov VB, Machs EM.** 2002. Cell cycle and its parameters in flowering plants. Tsiotologiia 44, 936–980.

**Hayashi K, Hasegawa J, Matsumaga S.** 2013. The boundary of the meiotic and elongation zones in roots: endoreduplication precedes rapid cell expansion. Scientific Reports 3, 2723.
Academy of Sciences, USA

tissue cultures: breakdown of normal controls. Proceedings of the National
Phillips RL, Kaeppler SM, Olhoft P.

plants. Springer-Verlag: Berlin, 207–239.

cell culture performance. In: Chandra S, ed. Biotechnology for medicinal

115–128.

1996. Cell synchronization and isolation of metaphase chromosomes from maize (Zea mays). root tips for flow cytometric analysis and sorting.

2009. The B73 maize genome: complexity, diversity, and dynamics. Science 326, 1112–1115.

Lee M, Phillips RL. 1988. The chromosomal basis of somaclonal variation. Annual Review of Plant Physiology and Plant Molecular Biology 39, 413–437.

Lee TJ, Pascuzzi PE, Settlage SB, et al. 2010. Arabidopsis thaliana chromosome 4 replicates in two phases that correlate with chromatid state. PLoS Genetics 6, e1000982.

Lee TJ, Shultz RW, Hanley-Bowdoin L, Thompson WF. 2004. Establishment of rapidly proliferating rice cell suspension culture and its characterization by fluorescence-activated cell sorting analysis. Plant Molecular Biology Reporter 22, 259–267.

Lucretti S, Nardi L, Nisini PT, Moretti F, Gualberti G, Dolezel J. 1999. Bivariate flow cytometry DNA/BrdUrd analysis of plant cell cycle. Methods in Cell Science 21, 155–166.

Matsutomo T, Wu J, Kanamori H, et al. 2005. The map-based sequence of the rice genome. Nature 436, 793–800.

Mayer KF, Waugh R, Brown JW, et al. 2002. Sequence of the rice genome. Nature 413, 697–703.

Menges M, Murray JA. 2002. Synchronous Arabidopsis suspension cultures for analysis of cell-cycle gene activity. The Plant Journal 30, 203–212.

Menges M, Samland AK, Planchas S, Murray JA. 2006. The D-type cyclin CYCD3;1 is limiting for the G1-to-S-phase transition in Arabidopsis. The Plant Cell 18, 893–906.

Muggeo VMR. 2008. Segmented: an R package to fit regression models with broken-line relationships. R News 8, 20–25.

Nagata T, Nemoto Y, Hasezawa S. 1992. Tobacco BY-2 cell line as the ‘HeLa’ cell in the cell biology of higher plants. International Review of Cytology 132, 1–30.

Oford KW, Scadden DT. 2008. Deconstructing stem cell self-renewal: genetic insights into cell-cycle regulation. Nature Reviews Genetics 9, 115–128.

Patil RA, Roberts C. 2013. Implications of cellular heterogeneity on plant cell culture performance. In: Chandra S, ed. Biotechnology for medicinal plants. Springer-Verlag: Berlin, 207–239.

Phillips RL, Kaeppler SM, Olhoft P. 1994. Genetic instability of plant tissue cultures: breakdown of normal controls. Proceedings of the National Academy of Sciences, USA 91, 5222–5226.