A journey through the microscopic ages of DNA replication

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Abstract Scientific discoveries and technological advancements are inseparable but not always take place in a coherent chronological manner. In the next, we will provide a seemingly unconnected and serendipitous series of scientific facts that, in the whole, converged to unveil DNA and its duplication. We will not cover here the many and fundamental contributions from microbial genetics and in vitro biochemistry. Rather, in this journey, we will emphasize the interplay between microscopy development culminating on super resolution fluorescence microscopy (i.e., nanoscopy) and digital image analysis and its impact on our understanding of DNA duplication. We will interlace the journey with landmark concepts and experiments that have brought the cellular DNA replication field to its present state.

Keywords DNA · DNA replication · Replication foci · Replicon · Replisome · Super resolution microscopy

The (very) early years

Long after water-filled glass bowls were used to read small letters (Singer 1914), a simple single lens microscope started the microscopic revolution (Bacon 1267) (see Table 1 and Fig. 1). Spurred throughout the ages by accidental inventions (van der Aa 1851), leaps by Galileo (Galilei 1610), and Hooke (1665), it was not until Carl Zeiss started to mass-produce microscopes in 1847 that DNA observation started to take off. Simultaneously, Mendel studied 29,000 pea plants (1866) and Haeckel postulated the containment of hereditary traits in the nucleus (1866) (Dahm 2008; Haeckel 1866), while Miescher put the microscope to good use and purified the nuclei and observed DNA (Miescher 1871). Köhler’s game-changing illumination technique (Köhler 1893) helped to perfect Zeiss UV-microscope together with Siedentopf in 1908. In 1927, shortly after Levene described the nucleic acid structure (1919), Koltsov postulated the semiconservative replication idea (Soyfer 2001).

The race down to the DNA structure and duplication

Phase contrast microscopy (Zernike 1955) and DNA X-ray diffraction images (Astbury 1947) Franklin, 1952, “Photo 52”) lead to fantastic images, new discoveries, and the description of the double helix DNA structure (Watson and Crick 1953). Meselson and Stahl ingeniously demonstrated the semiconservative mode of DNA replication (Meselson and Stahl 1958). The theoretical description of a confocal microscope and the first practical application of a Nipkow disk in microscopy (Egger and Petráň 1967; Petráň et al. 1968) were clear landmarks of the microscopy revolution.

Radioactive labeling and autoradiography allowed Cairns to observe DNA unwinding and the replication fork (Cairns 1963), and Huberman and Riggs observed similar replication structures in mammalian chromosomes (Huberman and Riggs 1966) and Okazaki described the lagging strand synthesis and “its” fragments (Okazaki et al. 1968; Okazaki and Okazaki 1969; Sugimoto et al. 1969; Sugimoto et al. 1968).
| Year | Landmark                                                                 | Author                      |
|------|--------------------------------------------------------------------------|-----------------------------|
| 63   | Water filed glass bowls to read small letters                           | (Singer 1914)              |
| 1267 | The first simple microscope                                             | (Bacon 1267)               |
| 1590 | Accidental discovery of the compound microscope with two (or more) lenses by Zacharias Janssen | (van der Aa 1851)          |
| 1610 | “Microscope” with ×1000 magnification                                   | (Galilei 1610)             |
| 1665 | “Micrographia”                                                           | (Hooke 1665; Singer 1914)  |
| 1847 | First “mass produced” microscopes in 1847                               | (Mendel 1866)              |
| 1866 | Hereditary traits in 29,000 pea plants                                   | (Dahm 2008; Haeckel 1866)  |
| 1871 | Purified nuclei for the first time and observed DNA                      | (Miescher 1871)            |
| 1893 | Ein neues Beleuchtungsverfahren für mikrophotographische Zwecke         | (Köhler 1893)              |
| 1907 | On the absorption of antibodies                                          | (Soyfer 2001)              |
| 1908 | First fluorescence microscopes based on UV-microscopy                    | (Knoll and Ruska 1932a)    |
| 1910 | Discovery of the electron microscope                                    | (Knoll and Ruska 1932b)    |
| 1947 | DNA X-ray diffraction images                                             | (Astbury 1947)             |
| 1953 | X-ray diffraction “Photo 51”                                             | (Watson and Crick 1953)    |
| 1953 | Discovery of the double-helix DNA structure                             | (Watson and Crick 1953)    |
| 1953 | Discovery of phase contrast microscopy                                   | (Zernike 1955)             |
| 1958 | Confirmation of the semiconservative DNA replication model              | (Meselson and Stahl 1958)  |
| 1957 | Discovery of the confocal microscope                                      | (Minsky 1961)              |
| 1962 | Extraction, purification, and properties of GFP                          | (Shimomura et al. 1962)    |
| 1963 | DNA unwinding for replication and “replication fork”                      | (Cairns 1963)              |
| 1966 | Autoradiography of chromosomal DNA fibers from Chinese hamster cells.   | (Huberman and Riggs 1966)  |
| 1966 | On the mechanism of DNA replication in mammalian chromosomes             | (Huberman and Riggs 1968)  |
| 1967 | First practical application of the “Nipkow disk” in confocal microscopy  | (Egger and Petráň 1967; Petráň et al. 1968) |
| 1968 | Mechanism of DNA chain growth. I. Possible discontinuity and unusual secondary structure of newly synthesized chains. | (Okazaki et al. 1968)      |
| 1968 | Mechanism of DNA chain growth. II. Accumulation of newly synthesized short chains in \( E. \) coli infected with ligase-defective T4 phages. | (Sugimoto et al. 1968)      |
| 1969 | Duration of the cell cycle                                               | (Van Dilla et al. 1969)    |
| 1969 | Mechanism of DNA chain growth. III. Equal annealing of T4 nascent short DNA chains with the separated complementary strands of the phage DNA | (Sugimoto et al. 1969)      |
| 1969 | Mechanism of DNA chain growth. IV. Direction of synthesis of T4 short DNA chains as revealed by exonucleolytic degradation. | (Okazaki and Okazaki 1969) |
| 1972 | Bidirectional Replication of Simian Virus 40 DNA                         | (Danna and Nathans 1972)   |
| 1974–1979 | Fork speed, replication speed, and replicon sizes                       | (Kriegerstein and Hogness 1974; Taylor 1977; Taylor and Hozier 1976; Wilson and Wilson 1975; Yurov 1977; Yurov 1978; Yurov 1979; Yurov and Liapunova 1977) |

1975 Continuous cultures of fused cells secreting antibody of predefined specificity.
1986 Structural organizations of replicon domains during DNA synthetic phase in the mammalian nucleus
1989 Three distinctive replication patterns
1992 Dynamic organization of DNA replication in mammalian cell nuclei spatially and temporally defined replication of chromosome
1992 Progression of DNA synthesis
1993 Structured Illumination Microscopy (SIM)
1994 Green fluorescent protein as a marker for gene expression
1994 4pi microscope
Table 1 (continued)

| Year | Landmark | Author |
|------|----------|--------|
| 1994 | Alignment and sensitive detection of DNA by a moving interface | (Bensimon et al. 1994) |
| 1997 | The replication origin decision point is a mitogen | (Wu and Gilbert 1997) |
| 1997 | Dynamic molecular combing: stretching the whole human genome for high-resolution studies. | (Michalet et al. 1997) |
| 1998 | Replicon clusters are stable units of chromosome structure evidence that nuclear organization contributes to the efficient activation and propagation of S phase in human cells | (Jackson and Pombo 1998) |
| 1999 | The spatial position and replication timing of chromosomal domains are both established in early G1 phase | (Dimitrova and Gilbert 1999) |
| 1999 | Single molecule analysis of DNA replication. | (Herrick and Bensimon 1999) |
| 2000 | Heterogeneity of eukaryotic replicons, replicon clusters, and replication foci | (Berezney et al. 2000) |
| 2000 | Dynamics of DNA replication factories in living cells | (Leonhardt et al. 2000) |
| 2000 | DNA replication at high resolution | (Keck and Berger 2000) |
| 2000 | Mechanisms of DNA replication | (Davey and O’Donnell 2000) |
| 2001 | Eukaryotic origins | |
| 2001 | Repression of origin assembly in metaphase depends on inhibition of RLF-BCdt1 by geminin | (Tada et al. 2001) |
| 2001 | Visualization of DNA replication on individual Epstein-Barr Virus episomes | (Norio and Schildkraut 2001) |
| 2002 | DNA polymerase clamp shows little turnover at established replication sites but sequential de novo assembly at adjacent origin clusters | (Sporber et al. 2002) |
| 2002 | DNA replication and chromatin | (Gerbi and Bielinsky 2002) |
| 2002 | Initiation of DNA replication in multicellular eukaryotes | (Gerbi et al. 2002) |
| 2003 | Sequence-independent DNA binding and replication initiation by the human origin recognition complex | (Vashee et al. 2003) |
| 2003 | The ‘ORC cycle’: a novel pathway for regulating eukaryotic DNA replication | (DePamphilis 2003) |
| 2004 | Stable chromosomal units determine the spatial and temporal organization of DNA replication | (Sadoni et al. 2004) |
| 2004 | DNA replication and DNA repair mechanisms most of the replication machinery is also used in DNA repair. | (Sancar and Lindsey-Boltz 2004) |
| 2005 | Preventing rereplication | (Blow and Dutta 2005) |
| 2005 | PCNA acts as a stationary loading platform for transiently interacting Okazaki fragment maturation proteins | (Sporber et al. 2005) |
| 2005 | Eukaryotic origins of DNA replication: could you please be more specific? | (Cvetic and Walter 2005) |
| 2006 | Origin selection and silent origins | (Patel et al. 2006) |
| 2006 | Regulating the licensing of DNA replication origins in metazoa | (DePamphilis et al. 2006) |
| 2006 | DNA replication: keep moving and don’t mind the gap. | (Langston and O’Donnell 2006) |
| 2007 | Impact of chromatin structure | |
| 2007 | Replisome mechanics: insights into a twin DNA polymerase machine. | (Pomerantz and O’Donnell 2007) |
| 2007 | The many faces of the origin recognition complex | (Sasaki and Gilbert 2007) |
| 2007 | High-throughput mapping of origins of replication in human cells. | (Lucas et al. 2007) |
| 2007 | Characterization of a triple DNA polymerase replisome. | (McInerney et al. 2007) |
| 2007 | Dynamic DNA helicase-DNA polymerase interactions assure processive replication fork movement. | (Hamdan et al. 2007) |
| 2007 | Polymerase switching in DNA replication. | (Lovett 2007) |
| 2008 | 3D–SIM | (Gustafsson et al. 2008) |
| 2008 | Division of labor at the eukaryotic replication fork. | (Nick McElhinny et al. 2008) |
| 2008 | DNA polymerases at the replication fork in eukaryotes | (Stillman 2008) |
| 2008 | Discovery of stimulated emission depletion (STED) | (Schmidt et al. 2008) |
| 2009 | In DNA replication, the early bird catches the worm. | (Boye and Grallert 2009) |
| 2009 | G-quadruplex structures: in vivo evidence and function. | (Lipps and Rhodes 2009) |
| 2009 | Eukaryotic DNA replication control: lock and load, then fire. | (Remus and Diffley 2009) |
| 2010 | Organization of DNA replication | (Chagin et al. 2010) |
| Year | Landmark                                                                 | Author                                                                 |
|------|--------------------------------------------------------------------------|----------------------------------------------------------------------|
| 2010 | Eukaryotic chromosome DNA replication: where, when, and how?             | (Masai et al. 2010)                                                   |
| 2010 | SCF (Cyclin F) controls centrosome homeostasis and mitotic fidelity through CP110 degradation. | (D’Angiolella et al. 2010)                                           |
| 2010 | Uncoupling of sister replisomes during eukaryotic DNA replication.      | (Yardimci et al. 2010)                                               |
| 2010 | DNA replication: making two forks from one prereplication complex.      | (Botchan and Berger 2010)                                            |
| 2011 | Eukaryotic origin-dependent DNA replication in vitro reveals sequential action of DDK and S-CDK kinases. | (Heller et al. 2011)                                                |
| 2011 | Failure of origin activation in response to fork stalling leads to chromosomal instability at fragile sites. | (Ozeri-Galai et al. 2011)                                           |
| 2011 | Selective bypass of a lagging strand roadblock by the eukaryotic replicative DNA helicase. | (Fu et al. 2011)                                                   |
| 2011 | Genome-wide depletion of replication initiation events in highly transcribed regions. | (Martin et al. 2011)                                               |
| 2011 | Origin association of Sld3, Sld7, and Cdc45 proteins is a key step for determination of origin-firing timing. | (Tanaka et al. 2011)                                               |
| 2012 | Genome-scale identification of active DNA replication origins.          | (Cayrou et al. 2012)                                                |
| 2012 | Forkhead transcription factors establish origin timing and long-range clustering in *S. cerevisiae* | (Knott et al. 2012)                                              |
| 2012 | A fragment based click chemistry approach towards hybrid G-quadruplex ligands: design, synthesis and biophysical evaluation | (Ritson and Moses 2012)                                           |
| 2012 | Histone hypoacetylation is required to maintain late replication timing of constitutive heterochromatin. | (Casas-Delucchi et al. 2012)                                      |
| 2012 | OriDB, the DNA replication origin database updated and extended.         | (Siow et al. 2012)                                                 |
| 2012 | Replication timing: the early bird catches the worm.                   | (Douglas and Diffley 2012)                                          |
| 2012 | CK2 inhibitor CX-4945 suppresses DNA repair response triggered by DNA-targeted anticancer drugs and augments efficacy: mechanistic rationale for drug combination therapy. | (Siddiqui-Jain et al. 2012)                                       |
| 2012 | Experimental approaches to identify cellular G-quadruplex structures and functions. | (Di Antonio et al. 2012)                                           |
| 2012 | Activation of the replicative DNA helicase: breaking up is hard to do.  | (Boos et al. 2012)                                                 |
| 2012 | Analysis of DNA replication profiles in budding yeast and mammalian cells using DNA combing. | (Bianco et al. 2012)                                              |
| 2012 | DeOri: a database of eukaryotic DNA replication origins.               | (Gao et al. 2012)                                                  |
| 2012 | Replication origins run (ultra) deep.                                  | (Gilbert 2012)                                                     |
| 2012 | Unraveling cell type-specific and reprogrammable human replication origin signatures associated with G-quadruplex consensus motifs. | (Besnard et al. 2012)                                             |
| 2012 | Targeted manipulation of heterochromatin rescues MeCP2 Rett mutants and re-establishes higher order chromatin organization. | (Casas-Delucchi et al. 2012)                                       |
| 2013 | Genome-wide mapping of human DNA-replication origins: levels of transcription at ORC1 sites regulate origin selection and replication timing. | (Dellino et al. 2013)                                             |
| 2013 | Functional implications of genome topology.                            | (Cavalli and Misteli 2013)                                         |
| 2013 | Nuclear positioning.                                                   | (Gundersen and Worman 2013)                                        |
| 2013 | Chromatin dynamics at the replication fork: there’s more to life than histones. | (Whitehouse and Smith 2013)                                       |
| 2013 | Quantitative, genome-wide analysis of eukaryotic replication initiation and termination. | (McGuffee et al. 2013)                                           |
| 2013 | The Elg1 replication factor C-like complex functions in PCNA unloading during DNA replication. | (Kubota et al. 2013)                                             |
| 2013 | Replication timing regulation of eukaryotic replicons: Rif1 as a global regulator of replication timing. | (Yamazaki et al. 2013)                                           |
| 2013 | Bubble-seq analysis of the human genome reveals distinct chromatin-mediated mechanisms for regulating early- and late-firing origins. | (Mesner et al. 2013)                                            |
| 2013 | A personal reflection on the replicon theory: from R1 plasmid to replication timing regulation in human cells. | (Masai 2013)                                                      |
| 2013 | From simple bacterial and archaeal replicons to replication N/U-domains. | (Hyrien et al. 2013)                                               |
| 2013 | Genomes and G-quadruplexes: for better or for worse.                   | (Tarsounas and Tijsterman 2013)                                    |
| 2013 | New insights into replication clamp unloading.                         | (Ulrich 2013)                                                     |
| 2013 | Replication dynamics: biases and robustness of DNA fiber analysis.     | (Técher et al. 2013)                                               |
Always look on the bright side

Along came *Aequorea victoria* green fluorescent protein (Shimomura et al. 1962) and brought light into darkness. Where audioradiography once ruled (Huberman and Riggs 1966; Huberman and Riggs 1968; Taylor et al. 1957), immuno-fluorescence labeling of fixed cells with monoclonal antibodies to modified nucleotides incorporated into newly synthesized DNA took the stage (e.g., Aten et al. 1992; Cardoso et al. 1993; Jackson and Pombo 1998; Jaunin et al. 1998; Ma et al. 1998; Mazzotti et al. 1990; Nakamura et al. 1986; Raska et al. 1989; Raska et al. 1991) only to be outshined by live cell microscopy of fluorescent fusion proteins (Cardoso et al. 1997; Leonhardt et al. 2000). Cell cycle duration (Van Dilla et al. 1969), fork speed, replication rate, and replicon sizes (Kriegstein and Hogness 1974; Taylor 1977; Taylor and Hozier 1976; Wilson and Wilson 1975; Yurov 1977; Yurov 1978; Yurov 1979; Yurov and Liapunova 1977) were all unearthed from the dark.

In parallel, the first affordable home computers made digital image analysis possible through the help of Wayne S. Rasband who developed the milestone in image analysis ImageJ (then, NIH Image) in 1987 (Schneider et al. 2012). Extensive microscopic analysis in fixed cells followed and provided a spatiotemporal description of replication sites (replication foci; see Fig. 2) in cells throughout S-phase (Nakamura et al. 1986) along with the three main distinctive early, mid, and late S-phase replication foci patterns (Jackson et al. 1962) and brought light into darkness. Where audioradiography once ruled (Huberman and Riggs 1966; Huberman and Riggs 1968; Taylor et al. 1957), immuno-fluorescence labeling of fixed cells with monoclonal antibodies to modified nucleotides incorporated into newly synthesized DNA took the stage (e.g., Aten et al. 1992; Cardoso et al. 1993; Jackson and Pombo 1998; Jaunin et al. 1998; Ma et al. 1998; Mazzotti et al. 1990; Nakamura et al. 1986; Raska et al. 1989; Raska et al. 1991) only to be outshined by live cell microscopy of fluorescent fusion proteins (Cardoso et al. 1997; Leonhardt et al. 2000). Cell cycle duration (Van Dilla et al. 1969), fork speed, replication rate, and replicon sizes (Kriegstein and Hogness 1974; Taylor 1977; Taylor and Hozier 1976; Wilson and Wilson 1975; Yurov 1977; Yurov 1978; Yurov 1979; Yurov and Liapunova 1977) were all unearthed from the dark.

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Fig. 1 Graphical overview of microscopy developments and their impact on DNA replication studies

Fig. 2 Organization of DNA replication from the genome to the individual replisome/replicon. A fluorescently labeled human HeLa Kyoto cell with a typical late S-phase replication pattern is presented in the top left corner (scale bar = 5 μm). Magnified super-resolution replication foci, with white circles representing individual replication sites displayed in the middle of the top row. A scheme of clustered DNA loops with active replication sites (white) is shown on the right. Starting point of DNA replication, the replication origin (ori), and the region replicated from a single origin displayed in the bottom row. Each replicon is replicated by two replication machineries (magenta), composed of various replication proteins, magnified in the bottom left corner. Adapted from (Chagin et al. 2016; Chagin et al. 2010)
and Pombo 1998; Mills et al. 1989; Nakayasu and Berezney 1989). Alongside, replication origins (Burhans et al. 1990; Burhans et al. 1991) were also reported.

The quest to replicate the genome

DNA loops (see Fig. 2) and their “functional” attachments to active transcription units were shown to be chromatin organizers during mitosis (Jackson et al. 1992), and replication factories were proposed as clusters of DNA replication sites organized by the nucleoskeleton (Hozák et al. 1993). Molecular combing, refined DNA fiber analysis, and sensitive detection of DNA (Bensimon et al. 1994) opened the door to whole genome stretching and high-resolution studies (Michalet et al. 1997). It allowed analysis of single DNA molecules undergoing replication (see Fig. 2) in a much greater resolution (Herrick and Bensimon 1999) than ever before. Stable replication clusters were also described as a basis for effective activation and propagation of genome replication during S-phase (Jackson and Pombo 1998) and regulation of replication timing (Dimitrova and Gilbert 1999).

Studies on DNA replication proteins (see Fig. 2) using live-cell fluorescence microscopy produced time lapse movies of replication factories and elucidated basic principles of their dynamic assembly-disassembly behavior (Leonhardt et al. 2000). Different regulatory levels were shown to be necessary to initiate and regulate DNA replication. Not only the chromatin structure, nuclear, and chromosomal locations but also origin recognition complex (ORC) and a whole bunch of other factors were found to define start sites of replication (DePamphilis 2003; Gerbi and Bielinsky 2002; Gerbi et al. 2002; Sasaki and Gilbert 2007).

Factories full of dominos

In addition to the “factory model” (Hozák et al. 1993), more dynamic models ensued (Sadoni et al. 2004; Sporbert et al. 2002) whereby replication at one site induces domino-like activation of neighboring origins, without the need to postulate pre-determined clusters of replicons. The combination with an earlier model postulating that origins of replication would be licensed only during mitosis and this license to replicate would be revoked after one round of replication (Blow and Dutta 2005; Blow and Laskey 1988) elegantly demonstrated how DNA is completely duplicated once, and only once, during each cell cycle. Despite Cvetic wishing for “eukaryotic origins of DNA replication to please be more specific” (Cvetic and Walter 2005), DNA replication origins in higher eukaryotes have been at best elusive. Nonetheless, as a whole, DNA replication is a very robust mechanism and stalled forks can be reactivated or reactivate neighboring origins to close all gaps and provide a perfect copy of billions of nucleotides at every cell division (Langston and O’Donnell 2006; Patel et al. 2006).

The ever elusive origin

The hunt for the elusive consensus motif of DNA replication origins continued with genome-wide high throughput mapping of potential origins and next-generation sequencing methods (Besnard et al. 2012; Cadoret et al. 2008; Cayrou et al. 2012; Dellino et al. 2013; Karnani et al. 2010; Lucas et al. 2007; Martin et al. 2011; Mesner et al. 2013; Mesner et al. 2011; Mukhopadhyay et al. 2014; Picard et al. 2014; Valenzuela et al. 2011) but stalled without a conclusive definition of the mammalian origin of replication. Correlations with specialized DNA structures (e.g., G-quadruplexes) and many others have been suggested but there seems not to be a simple solution and potentially there is no need to have one.

Studies into the epigenomic landscape, epigenetic control of DNA replication, and higher order chromatin organization (Casas-Delucchi and Cardoso 2011; Casas-Delucchi et al. 2012) have provided a link of epigenetic modifications (in particular, histone acetylation level) and temporal control of DNA replication origin firing.

Altogether, even Hyrien’s “Peaks cloaked in the mist,” all out approach was not able to identify possible origins by similarities in thousands of microarrays and/or next-generation sequencing data, suggesting origins form at unspecific DNA sites, but are suppressed by ongoing transcription (Hyrien 2015), which is highly correlated with histone acetylation.

To go where no one has gone before: beyond the Abbe limit

Meanwhile, the microscopy arms race to and beyond the diffraction limit calculated by Abbe continued with the Structured Illumination Microscopy (SIM) (Bailey et al. 1993), the 3D–SIM (Gustafsson et al. 2008) and the stimulated emission depletion (STED) (Schmidt et al. 2008).

The first attempts to label dating back to 1986 (Nakamura et al. 1986) and quantify replication sites in cells yielded numbers on the low hundreds (see Fig. 3). A decade later with the advent of digital imaging and computational image analysis tools, these numbers grew to around one thousand (Berezney et al. 1996; Fox et al. 1991; Jackson and Pombo 1998; Ma et al. 1998), where they remained for several years (see Fig. 3).

Such numbers of replication sites were compatible with a concept of clusters of replicons activated together and, thus, visualized together.

The dramatic increase in spatial resolution made possible with the new super-resolution microscopy techniques...
(fluorescence nanoscopy) enabled the visualization, for the first time, of smaller replication structures (Baddeley et al. 2010; Cseresnyes et al. 2009; Ligasová et al. 2009). It was now possible to resolve structures well below the Abbe limit, down to 30 nm and smaller. Nanoscopy (Gustafsson et al. 2008; Hell 2003; Hell et al. 1994) is in full swing and let us go where no one has gone before: beyond the Abbe limit. This, on the other hand, created another level of demand upon image analysis tools.

“Myths” confirmed!

The stage was now set to try and unveil the units of genome replication, i.e., the replicons and their associated machinery, the replisome, in cells.

From the earlier studies using light nanoscopy techniques (Baddeley et al. 2010; Cseresnyes et al. 2009) as well as electron microscopy (Koberna et al. 2005), suitable computational image analysis protocols were developed (Chagin et al. 2015). These combined efforts led to a further increase in the numbers of replication sites measured in cells (see Fig. 3), which was now finally compatible and fitting with the predicted numbers of replicons needed to duplicate the genome in human cells (Chagin et al. 2016; Lôb et al. 2016).

The microscopic information age had arrived. Previous efforts by Shaw et al. (2010), together with measurements throughout the years culminating on the visualization and quantification of individual replicons in cells in 4D, all supported by 3D–SIM imaging (Chagin et al. 2016) were all combined in a minimalistic but comprehensive 4D replicon simulation model (Lôb et al. 2016) displaying previously published replication polarity gradients, replication timing profiles, N/U domains, topologically associating domains, and timing transition regions (Audit et al. 2013; Baker et al. 2012; Chen et al. 2010; Hyrien et al. 2013; Pope et al. 2014).

Journey into the future

Future work should aim to bridge the ever-increasing genome-wide population data, with single molecule and single-cell microscopic data. Novel ways to combine and relate these very different types of information should be developed to get the highest spatial together with the highest temporal resolution without compromising the data on variability between single cells.

Importantly, the available models should be put to work to predict and test genome replication in different cell types and species and under different stress conditions. This would unleash the value of the existing models and lead us into the in silico DNA replication era.

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Conflict of interest The authors declare that they have no conflict of interest.
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