Zika Virus Disrupts Phospho-TBK1 Localization and Mitosis in Human Neuroepithelial Stem Cells and Radial Glia

Graphical Abstract

Highlights
- Derivation of human neocortical and spinal cord neuroepithelial stem (NES) cells
- Zika virus (ZIKV) infects NES cells and radial glia, impairing mitosis and survival
- ZIKV induces mitochondrial sequestration of centrosomal phospho-TBK1
- Nucleoside analogs inhibit ZIKV replication, protecting NES cells from cell death

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In Brief
Onorati et al. establish neuroepithelial stem (NES) cells as a model for studying human neurodevelopment and ZIKV-induced microcephaly. Together with analyses in human brain slices and microcephalic human fetal tissue, they find that ZIKV predominantly infects NES and radial glial cells, reveal a pivotal role for pTBK1, and find that nucleoside analogs inhibit ZIKV replication, protecting NES cells from cell death.

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Zika Virus Disrupts Phospho-TBK1 Localization and Mitosis in Human Neuroepithelial Stem Cells and Radial Glia

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SUMMARY

The mechanisms underlying Zika virus (ZIKV)-related microcephaly and other neurodevelopmental defects remain poorly understood. Here, we describe the derivation and characterization, including single-cell RNA-seq, of neocortical and spinal cord neuroepithelial stem (NES) cells to model early human neurodevelopment and ZIKV-related neuropathogenesis. By analyzing human NES cells, organotypic fetal brain slices, and a ZIKV-infected micrencephalic brain, we show that ZIKV infects both neocortical and spinal NES cells as well as their fetal homolog, radial glial cells (RGCs), causing disrupted mitoses, supernumerary centrosomes, structural disorganization, and cell death. ZIKV infection of NES cells and RGCs causes centrosomal depletion and mitochondrial sequestration of phospho-TBK1 during mitosis. We also found that nucleoside analogs inhibit ZIKV replication in NES cells, protecting them from ZIKV-induced pTBK1 relocalization and cell death. We established a model system of human neural stem cells to reveal cellular and molecular mechanisms underlying neurodevelopmental defects associated with ZIKV infection and its potential treatment.

INTRODUCTION

The human CNS is a complex organ that, as a result of its extended development, is susceptible to a host of genetic and environmental insults. Although great strides have been made in mapping the genetic landscape of human neurodevelopmental malformations, understanding of the mechanisms by which diverse environmental pathogens affect human neurodevelopment has been lagging (Bae et al., 2015; Diaz and Gleeson, 2009; Lui et al., 2011; Silbereis et al., 2016; Woods, 2004).

The emerging link between the mosquito-borne flavivirus Zika virus (ZIKV) infection of pregnant women and fetal microcephaly reinforces the need to use tissue- and species-relevant cellular systems to study human CNS development and establish experimental systems for modeling ZIKV infection, neurotropism, and treatment (Miner and Diamond, 2016; Schuler-Faccini et al., 2016). In adults, rare complications of ZIKV infection include Guillain-Barré syndrome (Broutet et al., 2016; Oehler et al., 2014) and meningoencephalitis (Araujo et al., 2016). More prominently, ZIKV infection during pregnancy is associated with, and likely causative for, severe fetal abnormalities including microcephaly/micrencephaly, lissencephaly, hydrocephaly, necrosis, periventricular and cortical calcifications, diffuse astrogliosis, hypoplasia of the brain stem and spinal cord (SC), Wallerian degeneration of the corticospinal tract, and ocular abnormalities (Brasil et al., 2016; Mlakar et al., 2016; de Paula Freitas et al., 2016; Rubin et al., 2016). More broadly, the classical teratogenic
TORCH syndrome pathogens (toxoplasma, other agents, rubella virus, cytomegalovirus, and herpes simplex virus) result in up to half of all perinatal deaths around the world, many associated with brain malformations including microcephaly, with an especially large burden in developing countries (Adams Waldorf and McAdams, 2013; Fine and Arndt, 1985).

Primary microcephaly results mainly from the depletion of neural stem/progenitor cells due to centrosomal defects, premature differentiation, and/or cell death (Diaz and Giese, 2009; Woods, 2004). Recently, ZIKV was shown to preferentially infect human pluripotent stem cell (hPSC)-derived neural progenitors and organoids and cause mitotic impairment and cell death in ZIKV mouse models (Dang et al., 2016; Garcez et al., 2016; Qian et al., 2016; Tang et al., 2016; Lazear et al., 2016; Li et al., 2016; Cugola et al., 2016; Miner et al., 2016; Wu et al., 2016). However, the human CNS is distinct in the diversity and proliferative potential of neural stem and progenitor cells (Lui et al., 2011; Bae et al., 2015; Silbereis et al., 2016; Gage and Temple, 2013). As a result, there may be aspects of viral infection that are unique to the human brain. Moreover, there have been only limited reports to date on human cell-type-specific responses to ZIKV over the course of infection primarily in in vitro or murine model systems without comparison to infected human brain tissue. Finally, it is not known to what extent microcephaly results from direct ZIKV infection of developing neural cells versus indirect effects, such as inflammation and altered placental support, which has been shown to affect brain development (Burton and Fowden, 2015; Mor, 2016). Addressing these questions in the context of the developing human CNS is crucial for deciphering ZIKV tropism and neuropathogenesis.

Here, we describe the derivation and characterization of neocortical (NCX) and SC neuroepithelial stem (NES) cells as in vitro models for neural stem/progenitor cells, early human neurodevelopment, and ZIKV-related neuropathogenesis. NES cell lines are derived from primary neural epithelial cells, the earliest population of resident neural stem cells present during neurodevelopment, when the neural tube is comprised of a pseudostratified neuroepithelium lining the central cavity (Bae et al., 2015). These cells constitute the ventricular zone (VZ) of the neural tube and serve as the stem cells of the CNS. Initially, neuroepithelial cells divide symmetrically in order to expand the stem tube and serve as the stem cells of the CNS. Moreover, after propagation in the presence of the apical basal junction cell lineages (aSVZ) and outer subventricular zone (oSVZ). These cell populations serve as the stem or progenitor cells for neurons and microglia (i.e., astrocytes and oligodendrocytes) and provide scaffolding for migrating nascent neurons (Bae et al., 2015). RGCs largely divide asymmetrically, giving rise to a daughter RGC, an intermediate progenitor cell (IPC), or a nascent neuron that subsequently migrates. Because of the ability to self-renew and differentiate, neuroepithelial cells are ideal candidates for the study of neural stem cell biology and various developmental diseases.

By comparing NES cells, organotypic fetal brain slices, and the postmortem human tissue in the context of ZIKV infection, we show that ZIKV infects both NCX- and SC-NES cells and their fetal homologs. RGCs in the VZ and SVZ, and to a lesser extent, postmigratory neurons in the cortical plate (CP; anlage of the cerebral cortical layers 2–6), ZIKV infection is associated with mitotic impairment, structural disorganization of the proliferative zones, and increased cell death of NES cells and RGCs. We also found that ZIKV causes subcellular relocation of the phosphorylated TANK binding kinase 1 (pTBK1), a pivotal player in antiviral innate immunity and cell proliferation (Farlik et al., 2012; Helgason et al., 2013; Pillai et al., 2015; Thurston et al., 2009) from centrosomes to mitochondria during mitosis of NES cells and RGCs. Antiviral nucleoside analogs, including the FDA-approved drug Sofosbuvir, inhibit ZIKV replication and ZIKV-induced cytopathology, including the relocation of pTBK1 to mitochondria, in NES cells. Together, these results validate the use of this NES cell system as an experimental model for studying early human neurodevelopment and ZIKV-induced pathogenesis, provide a testable hypothesis for the mechanism of ZIKV-induced cytopathology, and lay the groundwork for future research on ZIKV-related neurodevelopmental defects and therapy.

RESULTS

Derivation of Human Neocortical and Spinal Cord Neuroepithelial Stem Cells

Different early human neural stem/progenitor cell types, including neuroepithelial-like cells, have been derived from hPSCs and the developing human brain (Conti and Cattaneo, 2010; Edri et al., 2015; Elkabetz et al., 2008; Sun et al., 2008, Tailor et al., 2013). However, stable cell lines of neuroepithelial cells have not been derived from the dorsal forebrain (prospective NCX) or the SC. Thus, we attempted to generate stable long-term, self-renewing populations of NCX and SC neuroepithelial cells from ten human specimens ranging from 5 to 8 postconceptual weeks (pcw; Carnegie stage 15–17) (Figure 1A and Table S1).

Dissociated cells were plated and examined for the presence of neural rosettes reminiscent of the radial arrangement and apico-basal polarization established by neuroepithelial cells in the native neural tube (Figure 1B). In total, we derived and characterized five stable NCX and three stable SC neuroepithelial cell lines. Moreover, after propagation in the presence of the growth factors FGF2 and EGF as well as BDNF (Figures S1A and S1J), the neuroepithelial cell lines exhibited stem/progenitor cell characteristics and retained a high neurogenic capacity; thus, we termed them NCX- and SC-NES cells, respectively. Both NCX- and SC-NES cells were immunopositive for the neuroepithelial marker SOX1 and the pan-neuronal stem cell markers nestin, SOX2, and vimentin (VIM) (Figures 1C, 1D, S1B, and S1K). Metaphase NES cells were also observed and labeled with the anti-phospho-VIM antibody, and some displayed a typical basal process (Figure S1C).

Next, we investigated whether NES cells retained regional identity after long-term expansion. NCX-NES cells expressed FOXG1 and OTX2 (Figures 1E and S1D), key transcription factors demarcating proliferative zones of the early human forebrain (Onorati et al., 2014). On the other hand, SC-NES cells were positive for the caudal regional marker HOXB4 (Krumlauf et al., 1993) (Figure S1L), thus demonstrating maintenance of regional identity even after more than ten passages. These cell lines have
Figure 1. Derivation, Profiling, and ZIKV infection of NES Cells

(A) A schematic of experimental procedure. NES cells were derived from human developing neocortex (NCX) or spinal cord (SC). Donor-matched brain tissues and NCX-NES cells were subjected to single-cell RNA-seq. SC-NES cells are described in Figure S1.

(B) NCX-NES cells form rosette structures after 24 hr in culture. Scale bars represent 50 μm.

(C–E) NCX-NES cells are positive for nestin (NES), SOX2 (99.4% ± 0.8%, n = 973), SOX1 (87.8% ± 1.9%, n = 1265), CTNNB1, and FOXG1 (96.5% ± 1.1%, n = 1277). Scale bars represent 50 μm. Results are presented as mean ± SD.

(F) Violin plots showing expression of cell-type-specific marker genes in NCX-NES cells and donor-matched NCX tissue single cells. pan-NSC, pan-neural stem cell; vRGC, ventricular radial glia cell; oRGC outer radial glia cell; IPC, intermediate progenitor cell; Intern., interneuron; Ast., astrocyte; Olig., oligodendrocyte; Micro., microglia. y axis, log2 transformed RPKM.

(G) As in (F), violin plots of ventral forebrain marker genes in NCX-NES and donor-matched NCX single cells.

(H and I) t-SNE plots of all NES cell lines and brain single cells colored by origin of the cells (H) or by cell type clusters identified with SNN-Cliq clustering algorithm (I).

(J and K) Electron microscopy images of mock (J) or ZIKV-infected (K) NCX-NES cells. Nucleus (N) appears to be normal (J”), and cytoplasm is clear of ZIKV viral particles (J”) in mock condition. Nuclear fragmentation is evident in the ZIKV-infected condition. Viral particles are evident around nuclear envelope (red arrows in K”, K”). Scale bars represent 1 μm.

See also Figures S1–S4.
been expanded for more than 1 year and 35 passages with no evidence of chromosomal instability (Figures S1E and S1M).

To test the neurogenic potential of NES cells, we used an optimized differentiation protocol (Figure S1F). All NES cells gave rise to MAP2+ and TUBB3+ neurons and extended neurites (Figures S1G and S1H). We found that NCX-NES cells differentiated into neurons positive for BCL11B (also known as CTIP2), a marker of deep-layer excitatory projection neurons in the NCX (Figure S1H). SC-NES cells also gave rise to TUBB3-, MAP2-, and RBFOX3 (also known as NeuN)-positive neurons, whereas some exhibited a cholinergic phenotype, as expected for a SC neuronal population (Figures S1N and S1O). Furthermore, GFAP+ astroglial cells (Figures S1I and S1P) and occasional O4+ oligodendrocytes were detected (Figure S1Q), thus demonstrating the multipotential stem cell capacity of NES cells.

Human NES Cells Exhibit the Transcriptional Signature of Early Neural Stem Cells

To further characterize NCX-NES cells, we conducted single-cell RNA-seq on 439 expanded NCX-NES cells and 383 cells from 5–6 and 8 pcw donor-matched NCX (Tables S1 and S2); 362 and 255 cells, respectively, passed our quality-control pipeline (Supplemental Experimental Procedures). The majority of cells analyzed from the donor-matched NCX tissue samples expressed canonical marker genes for neuroepithelial cells and RGCs of the dorsal forebrain (Figure 1F). NCX-NES cells showed similar expression profiles as their donor-matched NCX cells (Figure 1F). Neither brain NCX cells nor NCX-NES cells expressed ventral forebrain markers (Figures 1G and S2D).

To place the transcriptional profiles of NCX-NES cells within the context of fetal development, we supplemented our donor-matched datasets with single-cell RNA-seq data from 526 cells collected from fetal NCX at 16, 19, and 20 pcw (Figure 1H); 392 of them passed quality control. We conducted principal component analysis for dimension reduction with genes as observations and single cells as variables (Figure S2A). The first principal component was used as the representation of each fetal time point or NCX-NES cell line. Pairwise Pearson’s correlation test showed that NCX-NES cells highly correlated with embryonic development time points (r > 0.8; Figure S2B). We next applied a clustering algorithm designed for single-cell studies, SNN-Cliq, to explore cellular composition over neocortical development and in NCX-NES cells. Based on the expression of marker genes for various cell types, we identified five NCX clusters (Figure S2C) representing ventricular and outer RGCs (vRGCs and oRGCs, respectively), two types of IPCs, and early neurons (Figure 1I). As expected from the expression profiles of NCX-NES cells, the largest of the three NCX-NES cell clusters highly correlated with the vRGC and IPC clusters, while the other two NCX-NES cell clusters showed neuroprogenitor-like gene expression profiles (Figure S2D). Together, these data establish NES cell lines as relevant models of early human CNS development.

Human NES Cells, but Not NES-Derived Neurons, Can Be Efficiently Infected by ZIKV

We sought to test whether the NCX-NES cell system can be used as a model for ZIKV infection. First, we infected NCX-NES cells with the 2010 Cambodian ZIKV strain FSS 13025 (MOI = 0.1), which shares 98% nucleotide identity with the current Brazilian epidemic strain. We monitored NCX-NES cells up to 6.5 days post-infection (DPI). At 3.5 DPI, we observed widespread ZIKV infection, as demonstrated by immunolabeling for the viral nonstructural protein 1 (NS1), with over 90% of NCX-NES cells infected (Figures S3A and S4F). We also observed a significant decrease in cell density over time compared to mock-infected cells (Figure S3B). Notably, the number of active caspase 3 (aCASP3+) apoptotic cells as well as pyknotic cells (i.e., cells showing condensation of the chromatin) was significantly higher in the infected group at 3.5 and 6.5 DPI compared to mock control (Figures S3C and S3D). Based on the analysis of immunolabeling for MKI67 (also known as Ki-67), a static marker of proliferative activity, we found that the proliferation levels were also significantly decreased by 3.5 DPI, with a continued decline through 6.5 DPI (Figures S3E and S3F). Electron microscopy also showed viral particles in infected NCX-NES cells and signs of nuclear fragmentation compared to mock cells (Figures 1J and 1K).

Recent studies have shown that hPSC-derived and mouse neural progenitors and neurons are infected by ZIKV, though neural stem and progenitor cells are preferential targets of infection (Qian et al., 2016; Tang et al., 2016). To test this in our model system, ZIKV was used to infect NCX-NES cells after differentiating them into neurons for 48 days. At 3.5 DPI, no obvious increase in the cytopathic effect or cell death was evident in the ZIKV-treated cultures (Figures S4A and S4B). In order to discriminate between ZIKV infection of neurons, which are postmitotic and express high levels of MAP2, and progenitors, we administered 5-bromo-2’-deoxyuridine (BrdU), a thymidine analog that incorporates into newly synthesized DNA, before ZIKV infection (Figure S4C). We found that only ~7.2% of the cells in the neuronal cultures were NS1+ (Figures S4D, S4E, and S4F). Among the NS1+ neurons, 46% were BrdU+, thus indicating that only ~3.3% of postmitotic neurons were infected by ZIKV (Figures S4G, S4H, and S4I). Therefore, the ability of ZIKV to infect neurons is extremely low in comparison to its ability to infect self-renewing NES cells (~90%, Figure S4F). This result also suggests that some NES cells infected during mitosis could have passed the infection onto daughter cells destined to become neurons.

ZIKV Preferentially Infects RGCs, Causing Scaffold Disorganization in Fetal Organotypic Slices

In addition to NES cells, we also infected ex vivo human organotypic fetal brain slices with ZIKV strain FSS 13025 (Figure 2A and Table S3). Before infection, brain slice cultures showed completely normal anatomical structures of the developing NCX, including an intact VZ/SVZ, intermediate zone/subplate (IZ/SP) and CP, with a prominent RG scaffold (Figure 2B). At 1.5 DPI, only a few cells were infected, and nearly all infected cells resided in the VZ/SVZ (Figure 2C); no obvious anatomical or cellular defects were otherwise observed. Interestingly, we did not observe any postmigratory neurons in the CP that were NS1+ at 1.5 DPI (Figure 2C). A dramatic increase in the number of NS1+ cells was evident at 3.5 and 6.5 DPI, mostly located in the VZ, SVZ, and IZ/SP (Figures 2F and 2G). A majority
of NS1-expressing cells also stained positive for the RGC marker VIM (Figures 2C, 2F, and 2G). Immunostaining of VIM showed that the RG scaffold was disorganized and that the radial organization of neurons in the CP was altered adjacent to highly infected regions of brain slices as compared to less infected regions or uninfected control (Figures 2D, 2E, 2F, and 2G). By 6.5 DPI, architectural disorganization was even more pronounced, with non-infected VIM+ RGCs acquiring a “balloon” shape, suggesting an indirect, non-cell-autonomous effect of ZIKV-infected cells (Figures 2F and 2G).

To test whether ZIKV-induced effects are virus strain specific, we tested the neurotropism of a Brazilian ZIKV strain, PE243, on NES cells, revealing an extensive NS1-positivity at 3.5 DPI (Figure S5A), which was not present in mock-infected cells or cells exposed to UV- or low pH-inactivated ZIKV particles. ZIKV PE243 also infected a neural primary culture from a 15 pcw NCX with a widespread NS1+ signal (Figure S5B). ZIKV PE243-infected organotypic brain slices also exhibited an architectural disorganization of the RG scaffold at 3.5 DPI. (Figure S5C).

Neocortical RGCs Contain ZIKV in an Infected Human Microcephalic Fetus

To confirm that ZIKV infection of RGCs is linked with microcephaly/microencephaly, we examined postmortem forebrain and SC tissues of the first reported ZIKV-infected microcephalic fetus (approximately 30 pcw, Table S4) from a ZIKV-infected mother (Mlakar et al., 2016) (Figure 3A). ZIKV RNA was previously found in the brain tissue of this fetus by RT-PCR (Mlakar et al., 2016). Tissue sections of the forebrain were immunopositive for NS1 and the ZIKV envelope (E) glycoprotein in the neocortical VZ and SVZ RGCs (Figures 3B–3E), which were identified by their position, morphology, and the expression of VIM (Figures 3G and 3I). Regions of the neocortical VZ/SVZ and RG morphology in those regions were disorganized in the infected brain, with numerous ventricular protrusions and a visually apparent decrease in the density of basal processes from RGCs as compared to an age-matched postmortem control (Figures 3F and 3H). RGCs also clustered abnormally in the VZ/SVZ, and the RG scaffold was disorganized, particularly in aCASP3+ cells, as indicated by nestin or VIM staining (Figures 3I, 3L, and 3M). Neurons in the NCX appeared disorganized and dysplastic, which could be due to either direct infection by ZIKV or indirectly through altered migration and neuron placement resulting from disruptions to the RG scaffold (Figure 3J). We did not observe infected cells in the ependymal layer, derivatives of RGCs lining the central canal of the SC (Figure 3K).

NES Cells and RGCs Express the Putative ZIKV Entry Receptors AXL and TYRO3

To better understand why ZIKV preferentially infects neural stem cells and RGCs, we surveyed the expression of putative ZIKV entry receptors in the NCX-NES cells and human fetal brain. Recent studies suggested that a number of cell-surface proteins may serve as ZIKV entry factors, including AXL, TYRO3, CD209, HAVCR1, and TIMD4, with AXL being the most plausible candidate (Hamel et al., 2015; Nowakowski et al., 2016). Among these five putative receptors, single-cell RNA-seq analysis found that AXL was generally expressed at the highest and most consistent levels in NCX-NES cells and RGCs and IPCs from the developing NCX (Figure 4A), whereas TYRO3 was expressed inconsistently and other receptors were essentially not expressed by NES cells, RGCs, and IPCs (Figures 4A and S2D). Analysis of previously published adult single-cell sequencing data (Darmanis et al., 2015) indicated that AXL was highly expressed in adult astrocytes, microglia, and endothelial cells but not in neurons (Figures 4B and S6A). Analysis of tissue-level transcriptome data from the developing human brain (Kang et al., 2011) showed that both AXL and TYRO3 expression in the NCX have bimodal patterns, with high expression in embryonic and early fetal samples and again increased expression beginning during late fetal development that remains at elevated levels after birth until adulthood (Figures 4C and 4D), which was consistent with the pattern of astrocyte generation (Zhang et al., 2016).

We corroborated the transcriptional data through immunofluorescence analysis of AXL protein expression and found co-expression of AXL and VIM in NCX-NES cells (Figures 4E and 4F). AXL expression in 14 pcw NCX was detected in VZ and VIM+ RGC basal processes as well as in blood vessels (Figure 4G). AXL was highly expressed in the VZ/SVZ but not the CP of fetal organotypic slice cultures (Figure 4H). Interestingly, NS1 staining was close to AXL staining in infected organotypic slices (Figure 4H). AXL signal was scarcely detectable in adult neocortical astrocytes, restricted to their end-feet contacting blood vessels, and in the periventricular astrocyte ribbon (Figure 4I). Taken together, these observations support AXL and possibly TYRO3 as viral entry factors and may explain the cell-type specificity of ZIKV infection in NES cells and RGCs (Nowakowski et al., 2016).

ZIKV Infection Redirects pTBK1 from Cytoplasm and Centrosomes to Mitochondria during Mitosis

Recent work reported that ZIKV disrupts mitosis and increases cell death in neural stem cells (Garcez et al., 2016; Qian et al., 2016), although the molecular basis of these effects is unknown. TANK-binding kinase 1 (TBK1) is a cellular serine-threonine kinase that is essential for both innate antiviral immune signaling and cell proliferation as well as other processes (Farlik et al., 2012; Helgason et al., 2013; Pillai et al., 2015; Thurston et al., 2009). We hypothesized that ZIKV could disrupt NES cell proliferation by diverting TBK1 activity away from mitosis and toward innate immune signaling. By using single-cell RNA-seq data, we found that TBK1 was highly expressed in both NES cells and RGCs/IPCs in the fetal NCX but not in immature neurons (Figure 5A). Single-cell RNA-seq data also showed that TBK1 is highly expressed in neurons, astrocytes, and microglia in the adult brain (Figure 5A). Consistent with this expression, analysis of a developing human brain transcriptome dataset (Kang et al., 2011) revealed a bimodal pattern of TBK1 expression in the NCX, with high expression in the embryonic and early fetal samples and again during postnatal development (Figure S6B).

To investigate whether TBK1 levels changed during ZIKV infection, we conducted droplet digital PCR (ddPCR) on both ZIKV-infected and mock-infected NES cells and organotypic brain slices. No differences in TBK1 levels were found between uninfected and ZIKV-infected (with either the Cambodian or
Figure 2. ZIKV Infects RGCs and Disrupts RG Scaffolding of Fetal Neocortical Organotypic Slices
(A) A schematic representation of the experimental procedure.
(B) Schematic illustration of the early fetal NCX. Vimentin (VIM) and DAPI immunostaining of pre-infection slices shows a normal architecture of neocortical wall. Higher magnification of CP and VZ reveals the radial organization of VIM+ RG fibers reaching the pial surface. MZ, marginal zone; CP, cortical plate; IZ/SP, intermediate zone/subplate; SVZ, subventricular zone; VZ, ventricular zone; PN, projection neuron; oRGC, outer radial glia cell; IPC, intermediate progenitor cell; vRGC, ventricular radial glia cell.
(C) Organotypic brain slices show ZIKV infection in RGCs at 1.5 days post infection (DPI); 80.6% ± 0.6% of the infected cells localized in VZ/SVZ, 19.4% ± 4% in the IZ/SP, and none in CP (total number of infected cells = 20). VZ/SVZ are significantly more infected than CP (paired t test, p < 0.05). The slices maintain a normal structure, with radial glial fibers reaching the pial surface and a normal orientation in VZ and SVZ.
(D and E) Non-infected neocortical slices show normal morphology and RG scaffold at both 3.5 (D) and 6.5 (E) DPI.
(F and G) ZIKV-infected slices show severe defects in RG scaffolding at 3.5 (F) and 6.5 (G) DPI. Most of the infected cells are located in VZ/SVZ and IZ/SP regions (73.3% ± 28.7% at 3.5 DPI; 89.3% ± 3.4% at 6.5 DPI), with fewer CP neurons showing infection (26.7% ± 28.7% at 3.5 DPI and 10.7% ± 3.4% at 6.5 DPI).
Brazilian ZIKV strains) NES cells or organotypic slices, though expression of TBK1 in the relatively homogeneous NES cell population was greater than in the organotypic brain slices, which are composed of both neural stem/progenitor cells and immature postmitotic neurons (Figure 5B). This indicates that ZIKV infection does not lead to obvious changes in TBK1 mRNA expression. (respectively; total number of infected cells = 456 at 3.5 DPI and 1,315 at 6.5 DPI). At 6.5 DPI, CP was significantly less infected than the other zones of the NCX (paired two-tailed t test, p < 0.01). Regions with high numbers of infected cells (shown in between arrowheads) show a more pronounced defect in radial orientation of RG fibers. Furthermore, it is possible to observe balloon-like cells that express VIM. Arrowheads indicate cells that are infected with ZIKV, immunopositive for NS1. Arrows show RG fibers. Scale bars represent 50 μm. Results are presented as mean ± SD. See also Figure S5.
Figure 4. NES Cells and RGCs Express Candidate ZIKV Entry Receptors AXL and TYRO3

(A and B) Violin plots of candidate entry factor gene expression distributions in NCX-NES and donor-matched NCX single cells (A) or adult NCX (B). y axis, log2 transformed RPKM (A) or log2 transformed RPM (B). Proj. neuro., projection neuron; Interneuro., interneuron; Oligo., oligodendrocyte; Endo. cells, endothelial cells.

(C and D) Expression trajectories of AXL (C) and TYRO3 (D) during human neurodevelopment in different brain regions. y axis represents gene expression quantified by log2 transformed signal intensities and the x axis represents the developmental ages quantified by the log10 transformed postconceptional days.

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expression in our two model systems. We therefore surveyed protein expression by immunoblotting for TBK1 and phosphorylated TBK1 (pTBK1) on human tissue from fetal VZ/SVZ and CP and found that there was a higher level of pTBK1 in VZ/SVZ compared to CP (Figure 5C). Furthermore, immunofluorescence analysis revealed that phospho-VIM\(^*\) mitotic NES cells co-expressed pTBK1 in centrosomes and midbody (Figure 5D). Consistent with this, pTBK1\(^*\) cells co-expressed PAX6 in the VZ/oSVZ of the fetal neocortical wall (Figure 5D). Together, these results suggest that ZIKV affects the localization or activity of TBK1 without substantially altering TBK1 expression.

We therefore examined the localization of pTBK1 in NCX-NES cells and organotypic slices during ZIKV infection. We found that pTBK1 localized to the centrosomes in NCX-NES cells and RGCs of organotypic brain slices, as demonstrated by co-staining of pTBK1 and TUBG (also known as γ-tubulin) (Figures 5E, 5G, 5K, and 5M). In contrast, pTBK1 was no longer detected in centrosomes in ZIKV-infected cells but was instead localized to mitochondria, as shown by colocalization with ATP5A1 (Figures 5F, 5H, 5L, 5N, and 5O). Relocalization of pTBK1 to mitochondria in ZIKV-infected NES cells was confirmed by immunoelectron microscopy (Figures 5I and 5J). ZIKV-infected cells also exhibited disrupted mitotic progression, including an increase in the number of NES cells and RGCs containing more than two centrosomes (Figures 5H, 5N, and 5P). Mitochondrial shape also appeared different, as shown by electron microscopy, with increased mitochondrial area in ZIKV-infected NES cells, whereas mitochondrial density was unchanged (Figure 5Q).

**TBK1 Inhibitors Impair Mitosis, Provoke Supernumerary Centrosomes in NES Cells, and Exacerbate ZIKV-Related NES Cell Death**

Given that the enrichment of pTBK1 at centrosomes of mitotic NES and RGCs is altered by ZIKV infection and that TBK1 has been functionally linked to mitosis (Pillai et al., 2015), we asked whether pharmacological inhibition of TBK1 kinase activity could mimic the impairment caused by ZIKV infection. We therefore treated NCX-NES cells with a small-molecule inhibitor of TBK1/IKK\(\alpha\), Amlexanox, which inhibits TBK1 activity but not its phosphorylation on Ser172 (Reilly et al., 2013). We found that Amlexanox caused a significant decrease in the number of pTBK1\(^*\) mitotic cells and increased the number of cells exhibiting supernumerary centrosomes 3.5 days after application (Figures 5R and 5S). Similar effects were demonstrated by another TBK1 inhibitor, BX795 (Figures 5R and 5S). These results are consistent with our previous finding that ZIKV-induced pTBK1 relocalization correlates with the neurodevelopmental pathology seen in the infected human brain.

TBK1 also instructs autophagy and apoptosis (Helgason et al., 2013). Therefore, inhibition of TBK1 may protect cells from cell death. We next investigated whether inhibiting TBK1 kinase activity impacts ZIKV-induced death of NCX-NES cells at 3.5 DPI. Application of Amlexanox or BX795 had no effect on cell death of uninfected cells (Figures S7A and S7B). However, both inhibitors modestly but significantly increased apoptotic cell death in ZIKV-infected NCX-NES cells (Figures S7A and S7B). Next, we also tested whether the two inhibitors can affect cell death in ZIKV-infected SC-NES cells at 3.5 DPI. Surprisingly, Amlexanox modestly decreased aCASP3\(^*\), whereas BX795 had no effect at 3.5 DPI (Figure S7C). We tested the effect of Amlexanox at 5.5 DPI and found that it increased aCASP3\(^*\) in SC-NES cells as observed in NCX-NES cells at 3.5 DPI (Figures S7E and S7F). Together, these results suggest that inhibition of TBK1 kinase activity can lead to severe cellular defects in mitotic NES cells and thus may exacerbate ZIKV-related cytopathology.

**Relocalization of pTBK1 by Other Viruses and Innate Immune Stimulation**

Given that TBK1 activation is an important component of innate antiviral signaling, we examined whether infection with another flavivirus, dengue-2 virus (DENV), or the TORCH pathogen, human cytomegalovirus (HCMV), could induce similar relocalization of pTBK1. Both DENV and HCMV infected NES cell cultures (Figures 6A and 6B) but had different effects on pTBK1 localization. Although HCMV also induced mitochondrial targeting of pTBK1, DENV did not (Figures 6A and 6B). This could be because DENV is known to inhibit the transduction of cytosolic RIG-I-like receptor (RLR) signaling through mitochondrial antiviral signaling (MAVS) and TBK1 to initiate cellular innate antiviral programs (Dalrymple, Cimica, and Mackow, 2015; He et al., 2016). Nevertheless, both DENV and HCMV induced cell death via CASP3 activation (Figure 6C). These data show that TBK1 activation and mitochondrial localization of pTBK1 is not specific to ZIKV.

To determine whether RLR activation alone could induce the relocalization of pTBK1 from centrosomes to mitochondria, we treated NES cells with KIN1408, a small-molecule agonist of RLRs (Pattabhi et al., 2016). Indeed, KIN1408 treatment induced relocalization of pTBK1 from centrosomes to mitochondria (Figures 6D and 6E) and increased CASP3-mediated cell death (Figures 6D and 6F). Together, these data suggest that ZIKV may
EGZIKV

Mock FH J

KM

LN

CP

A B

TBK1

pTBK1

GAPDH

ZIKV/SVZ

Fetal neocortex (16 pcw)

C

VZ/SVZ CP

TBK1

pTBK1

GAPDH

NES cells

Fetal neocortical organotypic slices

pTBK1 ATP5A1 NS1 DAPI pTBK1 TUBG NS1 DAPI pTBK1 immunoelectron microscopy

Mock

ZIKV

NES cells

Slice cultures

NES cells

Slice cultures

NES cells

Vehicle Amlexanox BX795

Vehicle Amlexanox BX795

Vehicle Amlexanox BX795

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activate RLRs, causing relocalization of pTBK1 from mitotic centrosomes to mitochondria, disruption of mitosis, and induction of cell death.

**Nucleoside and Nucleotide Analog Inhibit ZIKV Replication and Protect NES Cells**

There is an urgent need to identify therapeutic approaches for halting ZIKV infection, especially in pregnant women (Barrows et al., 2016). Chain-terminating nucleoside analogs have proven to be potent, broad-spectrum antivirals. Indeed, a few nucleoside analogs have already been shown to inhibit ZIKV in cell culture and mouse models of infection (Zmurko et al., 2016; Eyer et al., 2016), although the antiviral activity, pharmacology, and toxicity of these specific molecules in humans are unknown. We therefore examined whether nucleoside analogs that show potent inhibition of hepatitis C virus (HCV), another member of the Flaviviridae, could inhibit ZIKV infection in NES cell cultures. An initial screen of several HCV inhibitors in HuH-7 hepatoma cells revealed that the nucleoside analog 2′-C-methyladenosine and the nucleotidyl produg Sofosbuvir showed antiviral activity against ZIKV, whereas Mericitabine, Balapiravir, PSI-6206, and 4′-azidocytidine analogs HY-77651 and HY-77652, lacked ZIKV antiviral activity (data not shown). Importantly, 2′-C-methyladenosine and Sofosbuvir both had potent anti-ZIKV activity in NES cells (Figures 7A and 7B). Sofosbuvir protected NES cells from ZIKV-induced cell death, whereas 2′-C-methyladenosine exhibited mild toxicity on its own (Figures 7A and 7C). Furthermore, both compounds inhibited the relocalization of pTBK1 (Figure 7D). Together, these data suggest that Sofosbuvir, a drug approved for treatment of chronic HCV, shows promise in protecting neural stem cells from ZIKV infection and pathogenesis.

**DISCUSSION**

The increased incidence of microcephaly linked to maternal ZIKV infection has created an international public health crisis. Understanding ZIKV cell tropism, the impact of ZIKV on cell biology, and how these effects lead to microcephaly and other neurodevelopmental phenotypes requires either direct access to developmental human brain tissue or the development of suitable model systems. In this study, we present several key advances. First, we report on the derivation and characterization of NES cells, a model system of long-term neuroepithelial stem cells derived from the NCX or SC. We also show that NES cells and RGCs are infected, mitotically impaired, and killed by ZIKV. We describe the relocation and sequestration of pTBK1 from centrosomes to mitochondria following ZIKV infection. Furthermore, we show that inhibition of TBK1 phenocopies ZIKV-infection-associated mitotic defects and supernumerary centrosomes. Finally, we found that two nucleoside analogs, one of which is an approved drug, can inhibit ZIKV replication...
and cell death in NES cells. These results are consistent with and substantially add to recent studies on ZIKV neurotropism and potential entry receptors (Miner and Diamond, 2016).

Several neurogenic populations of stem/progenitor cells have been developed from mouse or human neural tissue, but many of these culture systems have limited neurogenic and differentiation potentials (Onorati et al., 2011; Sun et al., 2008). Although recent work on hPSCs has expanded the range of neural cell types attainable (Conti and Cattaneo, 2010; Edri et al., 2015; Koch et al., 2009) and a population of NES cells has been derived from the hindbrain (Tailor et al., 2013), the NES cell system we described and characterized is, to our knowledge, the first neuropotent populations derived directly from the prospective NCX and SC, respectively. As such, NES cells represent a substantial advance facilitating the study of early development and neuro-

By comparing our NES cells to ex vivo fetal organotypic brain slices and a ZIKV-infected fetal brain, we demonstrated definitively that RGCs are the primary cell type infected by ZIKV in the prenatal human brain, as previously suggested by using hPSC-derived neural progenitors or midfetal stem cell systems (Dang et al., 2016; Garcez et al., 2016; Qian et al., 2016; Tang et al., 2016; Lazear et al., 2016; Li et al., 2016; Cugola et al., 2016; Miner et al., 2016; Wu et al., 2016; Hanners et al., 2016; http://www.biorxiv.org/content/biorxiv/early/2016/06/15/058883.1.full.pdf). We also found that early SC stem cells can be infected by ZIKV. This is in agreement with a previous report showing SC defects in ZIKV-infected mouse models (Lazear et al., 2016) as well as a human case where descending tracts of the SC, particularly the lateral corticospinal tracts, were very affected (Mlakar et al., 2016), though this latter phenotype could be a secondary pathology in the human CNS. Additionally, by conducting a comprehensive single-cell RNA-seq analysis of NES cells and human prenatal NCX specimens, including matched donor tissue, we have also generated a resource for future neuroscience and genomic studies.
consequence of neocortical infection. Why ZIKV infection is associated with micrencephaly rather than SC defects may in part be a result of the timing at which infection happens; human SC completes neurogenesis much earlier than the rest of the CNS (Silbereis et al., 2016), and so if infection occurs after the end of the first trimester, then SC cells may rarely be infected. However, the low percentage of neurons infected in NES cells and the fact that infected neurons in the organotypic slice cultures appeared only after the infection of VZ/SVZ RGCs suggest that some neurons may inherit the virus from infected RGCs.

Figure 7. Nucleoside and Nucleotide Analogs Inhibit ZIKV Replication and Protect NES Cells
(A) Representative pictures showing 2’-C-methyladenosine and Sofosbuvir treatment on ZIKV-infected NES cells at 3.5 DPI. At different concentrations, both drugs show a potent anti-ZIKV activity, as demonstrated by NS1 staining. pTBK1 localization is centrosomal in infected drug-treated cells. aCASP3 staining shows apoptotic cell death rate in the different conditions.
(B) Bar plot indicating the percentage of NS1+ cells in ZIKV-infected NES cells after vehicle or drug treatment at 3.5 DPI (number of NS1+ cells = 2,961 out of 3,501 total cells for vehicle, 7 out of 2,118 total cells for 2’-C-methyladenosine [20 µM], 2 out of 1,959 total cells for 2’-C-methyladenosine [40 µM], 136 out of 2,248 total cells for Sofosbuvir [20 µM], and 25 out of 2,099 total cells for Sofosbuvir [100 µM]). Two-tailed t test, *p < 0.05 and ***p < 0.001. n.s., not significant.
(C) Bar plot of the percentage of aCASP3+ cells in ZIKV-infected NES cells after vehicle or drug treatment at 3.5 DPI (number of aCASP3+ cells = 27 out of 1,403 total cells for vehicle, 14 out of 9,55 total cells for 2’-C-methyladenosine [20 µM]; 24 out of 502 total cells for 2’-C-methyladenosine [40 µM], 5 out of 657 total cells for Sofosbuvir [20 µM], and 4 out of 727 total cells for Sofosbuvir [100 µM]). Two-tailed t test, *p < 0.05 and **p < 0.01. n.s., not significant.
(D) Bar plot indicating the percentage of pTBK1+ cells that were localized in centrosomes or mitochondria in ZIKV-infected NES cells after vehicle or drug treatment at 3.5 DPI (number of pTBK1+ cells = 32 out of 2,098 total cells for vehicle, 5 out of 1,163 total cells for 2’-C-methyladenosine [20 µM], 4 out of 1,457 cells for 2’-C-methyladenosine [40 µM], 13 out of 1,591 cells for Sofosbuvir [20 µM], and 17 out of 1,373 total cells for Sofosbuvir [100 µM]). Fisher’s exact test, ***p < 0.001. Scale bars represent 20 µm.
Results are presented as mean ± SD.
We found that ZIKV infection of NES cells and RGCs caused a subcellular relocation of pTBK1 from centrosomal and cytosolic to mitochondrial locations. Consistent with its role in mitotic proliferation, pTBK1 was present in mitotic uninfected RGCs and NES cells throughout the cytoplasm and enriched at the centrosomes and midbody during cytokinesis. In contrast, we found that pTBK1 localized to mitochondria in ZIKV infected cells and was depleted in the cytoplasm and centrosomes. This is in agreement with the reported recruitment of TBK1 to the mitochondrial antiviral signaling platform in response to viral infection (Belgnaoui et al., 2011; Ning et al., 2014). Taken together, these results suggest a cell defense mechanism employed by NES cells and RGCs upon ZIKV infection that may result in abnormal mitoses. Although mitotic impairment in ZIKV infection has been described in previous reports (Garcez et al., 2016; Qian et al., 2016; Tang et al., 2016; Li et al., 2016), neither TBK1-associated centrosomal nor mitotic defects have been previously described.

Because altered cell-cycle dynamics may in some instances cause microcephaly, the removal of pTBK1 from the centrosome may explain some of the phenotypes associated with ZIKV infection. To test this hypothesis, we inhibited TBK1 by Amlexanox or BX795. Application of these inhibitors to NES cells impaired mitosis and induced supernumerary centrosomes. Multiple centrosomes and nuclei were also present in ZIKV-infected NES cells and RGCs, confirming that pTBK1 plays an important role in mitosis in these cells. Intriguingly, the majority of human microcephaly-associated genes encode centrosomal proteins (Diaz and Gleeson, 2009; Roberts et al., 2000; Woods, 2004). Moreover, studies in mutant mice and humans with mutations in centrosomal genes exhibit deficits including supernumerary centrosomes and failed cytokinesis, resulting in severely defective neurogenesis, massive apoptosis in the proliferative zones, and microcephaly (Bae et al., 2015; Diaz and Gleeson, 2009; Roberts et al., 2000; Woods, 2004).

pTBK1 is recruited to mitochondria in response to RNA virus infections, where it serves to transduce signals from MAVS protein and activate IRF-3 (Liu et al., 2015). Given that other flaviviruses induce RLR-mediated MAVS signaling (Lazar et al., 2013; Suthar et al., 2013), it is likely that mitochondrial relocalization of pTBK1 in ZIKV-infected NES cells and RGCs occurs via MAVS signaling. Supporting this possibility, treatment with KIN1408, a known RLR agonist, phenocopied both the relocalization of pTBK1 and the induction of CASP3-mediated cell death in NES cells. These data therefore support the model that ZIKV disrupts mitosis in neural stem cells by activating RLRs. Interestingly, the human TORCH pathogen, HCMV, caused a similar relocalization of pTBK1 and CASP3-mediated cell death. However, DENV, a flavivirus that inhibits RLR/MAVS/pTBK1 signaling (Darlymple, Cicimica and Mackow, 2015; He et al., 2016), did not induce pTBK1 relocalization while still resulting in apoptotic cell death. It is unlikely that the mitochondrial localization of pTBK1 alone is sufficient to explain the extensive cell death observed in more severe examples of ZIKV-infected fetal brain. The presence of uninfected RGCs exhibiting atypical cytomegalic “balloon” morphologies suggests that even uninfected neural cells can be non-cell-autonomously affected in response to ZIKV infection (Bell et al., 1971). Different cell types in the adult brain, including astrocytes, may be infected with ZIKV and nonetheless seem to survive for extended periods of time. In this and earlier studies in mouse models of ZIKV infection, neurons maintained normal morphology, despite being infected by ZIKV and likely survived for extended period of time. Similarly, we observed numerous RGCs and ependymal cells in ZIKV-infected postmortem human fetal NCX and SC that exhibited no obvious signs indicative of impending cell death. Crucially, these cells are either slowly dividing as neurogenesis has already ceased or are largely postmitotic, which raises the possibility that the process of mitosis, or attempting to go through mitosis in the absence of centrosomal pTBK1, may be the central mechanism underlying ZIKV-induced depletion of the neural progenitor pool and the subsequent microcephalic phenotype. Additionally, ZIKV may not affect neuronal migration even though the centrosome is critical for this process (Bae et al., 2015; Diaz and Gleeson, 2009; Woods, 2004).

We identified two nucleoside analogs, including the FDA-approved drug Sofosbuvir, that inhibit ZIKV infection and cytopathology in NES cells. Additional study is warranted to examine the efficacy of Sofosbuvir in inhibiting ZIKV in humans and test safety in pregnant women. In contrast, the TBK1 inhibitor Amlexanox, which is used clinically (Reilly et al., 2013), may exacerbate ZIKV-induced cytopathic effects.

Finally, our work may have significant implications concerning the effects of infection, the innate immune response, and the development of the CNS. The human CNS is usually considered an immunoprivileged organ and is generally protected from pathogens circulating in the rest of human body during development by the placenta (Burton and Fowden, 2015; Mor, 2016) and the forming blood-brain and blood-cerebrospinal fluid barriers (Marín-Padilla, 2012; Molgård and Saunders, 1980). However, some pathogens, such as ZIKV, possess the ability to cross the placenta and blood-brain barriers and infect the brain. Notably, almost all congenital TORCH infections involve some aspect of neurodevelopmental defect, including retinopathy, brain calcification, and microcephaly (Fine and Arndt, 1985; Neu et al., 2015). Considering shared clinical features and the pivotal role of TBK1 in the innate immune response to viruses and in autophagic clearance of various types of pathogens (Helgason et al., 2013; Thurston et al., 2009), we found that HCMV congenital infections may disrupt the development of human CNS through overlapping mechanisms that involve altering the subcellular localization and function of pTBK1 in human neural stem cells.

Future studies are expected to dissect the role of pTBK1 in innate immune response and the exact mechanisms by which centrosomal depletion of pTBK1 is triggered in the context of RLRs/MAVS signaling. Furthermore, the functional characterization of downstream effects of pTBK1 mitochondrial activation will reveal its role in cell death and potential molecular targets in a broad scenario of viral infection.

In summary, our study shows the utility of different in vitro and ex vivo model systems of human neural stem cells for studying early neurodevelopment and ZIKV infections and reveals cellular and molecular mechanisms underlying neurodevelopmental defects associated with ZIKV infection and, potentially, other pathogens.
EXPERIMENTAL PROCEDURES

ETHICS STATEMENT

This study was conducted using de-identified postmortem human specimens from tissue collection at the Department of Neuroscience, Yale School of Medicine, and tissue specimens provided by the Human Development Biology Resource (www.hdbr.org) and by the Laboratory of Developmental Biology at the University of Washington. Appropriate informed consent was obtained, and all available non-identifying information was recorded for each specimen. All work was performed according to NIH guidelines for the acquisition and distribution of human tissue for bio-medical research purposes and with approval by the Human Investigation Committees and Institutional Ethics Committees of each institute from which samples were obtained. Appropriate written informed consent was obtained, and all available non-identifying information was recorded for each sample. Details of the specimens analyzed are provided in the Supplemental Information.

HUMAN NES CELLS DERIVATION

NES cells were derived from dorsal forebrain and SC from de-identified prenatal human specimens. In brief, tissues were dissociated until a single-cell suspension was formed. Cells were resuspended and plated into coated dishes. See the Supplemental Experimental Procedures for details.

SINGLE-CELL RNA-SEQ ANALYSIS

Cells were captured using C1 Single-Cell Auto Prep Integrated Fluidic Circuit (Fluidigm). Single-cell sequencing libraries were generated according to manufacturer’s protocol. Reads were aligned using STAR (v2.4.0). Gene expression was calculated using RSEQTools and SAMtools. See the Supplemental Experimental Procedures for details and analysis pipelines.

EX VIVO FETAL ORGANOPTHIC SLICES CULTURES

Neocortical walls from four fetal brains were dissected in hibernate solution. Tissue was then imbedded in agarose and sectioned. 300 μm slices were transferred to cell culture with neuronal medium. See the Supplemental Experimental Procedures for details.

PREPARATION OF VIRUS STOCKS

ZIKV strain F5S 13025 was obtained from the World Reference Center for Emerging Viruses and Arboviruses at University of Texas Medical Branch, Galveston. The Brazilian ZIKV strain PE243 was recovered from a 19-year-old female in Brazil in late 2015. DENV-2 strain 16681 was recovered from the cDNA clone pD2/IC-30P. See the Supplemental Experimental Procedures for details. Independent stocks of HCMV-GFP were kindly supplied by Felicia Goodrum and Scott Terhune.

ZIKV INFECTION OF NES CELLS AND FETAL ORGANOPTHIC BRAIN SLICES

For NES cells, cells were seeded 1 day before virus addition. All viral infections were performed under the same conditions at an MOI of 0.1 for 2 h followed by three washes and addition of fresh NES medium. For fetal organotypic brain slices, viruses were added to culture medium. See the Supplemental Experimental Procedures for details.

APPLICATION OF SMALL-MOLECULE INHIBITORS

Amlexanox was applied at 10 μM, BX795 at 1 μM, and KIN1408 at 10 μM (final concentration). Sofosbuvir and 2’-C-methyladenosine were applied at the indicated concentrations. See the Supplemental Experimental Procedures for details.

BrdU LABELING, IMMUNOSTAINING, AND ELECTRON MICROSCOPY

10 μM BrdU was administered 30 min prior to viral infection and then continued until cell harvesting. For immunostaining, fixed samples were permeabilized and blocked at room temperature, incubated with primary antibodies overnight at 4°C, washed and incubated with secondary antibodies. All images were acquired using a fluorescent microscope or a laser scanning confocal microscope. For detailed methods and electron microscopy, see the Supplemental Experimental Procedures.

RNA ISOLATION, DROPLET DIGITAL PCR, AND IMMUNOBLOTTING

Total RNA or proteins were extracted from mock and infected samples as described in Supplemental Experimental Procedures.

STATISTICAL ANALYSIS

The presented data are mean ± SD or SEM from three separate experiments. Differences between the groups were analyzed using a Student’s t test. Values of all significant correlations are given with degree of significance indicated (* p < 0.05, ** p < 0.01, and *** p < 0.001). For organotypic slice culture, quantification was performed in up to four sections that were representative of at least three slices. Cells were counted as described in the respective figure legend. The number of cells with pTBK1 located in centrosomes or mitochondria follows a binomial distribution. A Fisher’s exact test was performed in order to compare the control and the infected and treated group. Similarly, to test whether infection changed the number of centrosomes, the cells were binned by the number of centrosomes (1–5), and a Fisher’s exact test was performed in order to compare control with infected and treated cells.

ACCESSION NUMBERS

The accession number for single-cell RNA-seq data reported in this paper is NCBI GEO: GSE81475. The accession number for the Brazilian ZIKV strain PE243 is GenBank: KX197192.

SUPPLEMENTAL INFORMATION

Supplemental Information contains Supplemental Experimental Procedures, seven figures, and four tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2016.08.038.

AUTHOR CONTRIBUTIONS


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