Quantitative live-cell imaging yields novel insight into endogenous WNT/CTNNB1 signaling dynamics

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Abstract

WNT/CTNNB1 signaling regulates tissue development and homeostasis in all multicellular animals. Multiple aspects of the underlying molecular mechanism remain poorly understood and critical information on endogenous WNT/CTNNB1 signaling dynamics is missing. Here we combine CRISPR/Cas9-mediated genome editing and quantitative live-cell microscopy to measure diffusion characteristics of fluorescently tagged, endogenous CTNNB1 in human cells with high spatiotemporal resolution. State-of-the-art functional imaging reveals that both in the absence and presence of WNT, a substantial fraction of CTNNB1 resides in slow-diffusing complexes in the cytoplasm and that WNT stimulation changes their identity. We also measure the concentration of complexed and free CTNNB1 in both the cytoplasm and the nucleus before and after WNT stimulation, and use these parameters to build a minimal computational model of WNT/CTNNB1 signaling. Our work reveals that WNT regulates the dynamic distribution of CTNNB1 across different functional pools by modulating the destruction complex, nucleocytoplasmic shuttling and nuclear retention.
Introduction

WNT signaling is one of the most ancient pattern-forming cell signaling cascades. It drives many biological processes from the onset of embryogenesis until adulthood in all multicellular animals (reviewed in van Amerongen and Nusse, 2009; Holstein, 2012; Loh et al., 2016). WNT signaling remains important throughout the lifespan of the organism and controls stem cell maintenance in many mammalian tissues, including the breast, intestine and skin (Barker et al., 2007; Lim et al., 2013; van Amerongen, Bowman, & Nusse, 2012). Disruption of the pathway causes disease, with hyperactivation being a frequent event in human colorectal and other cancers (reviewed in Nusse and Clevers, 2017; Polakis, 2000; Wiese et al., 2018; Zhan et al., 2017).

The key regulatory event in WNT/CTNNB1 signaling (traditionally known as ‘canonical WNT signaling’) is the accumulation and nuclear translocation of the transcriptional co-activator beta-catenin (CTNNB1). In the absence of WNT signaling, rapid turnover by the so-called destruction complex maintains low levels of CTNNB1 (Figure 1A). The destruction complex consists of the scaffold proteins APC and AXIN, which bind CTNNB1, and the serine/threonine kinases CSNK1 and GSK3, which subsequently phosphorylate residues S45, T41, S37 and S33 (Amit et al., 2002; C. Liu et al., 2002). This primes CTNNB1 for ubiquitination by E3 Ubiquitin Protein Ligase BTRC and subsequent proteasomal degradation (Aberle, Bauer, Stappert, Kispert, & Kemler, 1997; Latres, Chiaur, & Pagano, 1999). In the current working model for WNT/CTNNB1 signaling, binding of WNT ligands to the FZD/LRP receptor complex sequesters and inhibits the destruction complex at the membrane in a process that involves DVL (Bilic et al., 2007; Schwarz-Romond et al., 2007). This allows newly synthesized CTNNB1 to accumulate (Figure 1B). Upon stimulation, CTNNB1 also translocates to the nucleus, where CTNNB1 binds to TCF/LEF transcription factors to regulate target gene transcription as part of a larger
transcriptional complex (Behrens et al., 1996; Fiedler et al., 2015; Molenaar et al., 1996; van Tienen, Mieszczanek, Fiedler, Rutherford, & Bienz, 2017).

The working model for WNT/CTNNB1 signaling described above, is the result of almost 40 years of research. The use of traditional genetic and biochemical approaches has allowed identification of the core players, as well as dissection of the main signaling events. However, multiple aspects of WNT/CTNNB1 signaling remain poorly understood. For instance the exact molecular composition and mechanism for inhibition of the destruction complex remain unclear (reviewed in Tortelote et al., 2017), and how WNT/CTNNB1 signaling regulates the subcellular distribution of CTNNB1 requires further scrutiny.

Most biochemical techniques lead to loss of spatial information and averaging of cell-to-cell heterogeneity, since proteins are extracted from their cellular context. Additionally, temporal information is usually limited to intervals of several minutes or hours. Live-cell microscopy offers better spatiotemporal resolution. However, currently many of these studies are conducted by overexpressing the protein(s) of interest. This can severely affect activation, localization and complex formation (T. J. Gibson, Seiler, & Veitia, 2013; Mahen et al., 2014).

Although stabilization of CTNNB1 by WNT signaling has been extensively studied, there are very few studies on the spatiotemporal dynamics of this process especially at the endogenous level (Chhabra, Liu, Goh, Kong, & Warmflash, 2019; Massey et al., 2019; Rim, Kinney, & Nusse, 2020).
Figure 1: Current model of the WNT/CTNNB1 pathway. A) In the absence of WNT ligands, free cytoplasmic CTNNB1 is captured by the destruction complex (or “degradosome”) consisting of AXIN, APC, CSNK1 and GSK3, where it is sequentially phosphorylated by the latter two kinases. This phosphorylation triggers ubiquitination by BTRC and subsequent proteasomal degradation. As a result, levels of CTNNB1 are kept low in both the cytoplasm and the nucleus. B) Binding of the WNT protein to the FZD and LRP receptors inhibits the destruction complex. This process is mediated by DVL and is thought to sequester the destruction complex to the membrane (also known as the “signalosome”). CTNNB1 accumulates in the cytoplasm and subsequently translocates to the nucleus, where it promotes the transcription of target genes, such as AXIN2, as a co-activator of TCF/LEF transcription factors. This transcriptional complex contains multiple other partners and is also termed the “enhanceosome”. Note that CTNNB1 also plays a structural role at the membrane in adherens junctions (not depicted).

Here we use CRISPR/Cas9 mediated genome editing in haploid cells to generate clonal cell lines that express fluorescently tagged CTNNB1. Using confocal imaging and automated cell segmentation we quantify the dynamic subcellular increase of endogenous CTNNB1 upon WNT stimulation. Moreover, using Fluorescence Correlation Spectroscopy (FCS) and Number and Brightness (N&B) analysis we measure the mobility and concentration of CTNNB1, providing information on CTNNB1 containing complexes in the cytoplasm and nucleus. Finally, we use these parameters to construct a novel computational model of WNT/CTNNB1 signaling. Our combined approach offers new insight into the dynamic regulation of CTNNB1 in mammalian cells through modulation of the destruction complex as well as nucleocytoplasmic shuttling and nuclear retention.
Results

Generation of clonal HAP1<sup>SGFP2-CTNNB1</sup> cell lines

To be able to visualize and quantify the spatiotemporal dynamics of WNT/CTNNB1 signaling at the endogenous level, we fluorescently tagged CTNNB1 in mammalian cells using CRISPR/Cas9 mediated homology directed repair (Ran et al., 2013) (Figure 2). To preserve the existing (epi)genetic control mechanisms of CTNNB1 expression, only the coding sequence for SGFP2, a monomeric, bright and photostable green fluorescent protein (G.-J. Kremers, Goedhart, van den Heuvel, Gerritsen, & Gadella, 2007), was seamlessly inserted at the starting ATG of the CTNNB1 coding sequence in HAP1 cells (Figure 2A, Figure 2 supplement 2A). HAP1 is a near-haploid and WNT responsive cell line (Andersson et al., 1987; Carette et al., 2011; Kotecki, Reddy, & Cochran, 1999; Lebensohn et al., 2016). By using haploid cells, we could ensure homozygous tagging of CTNNB1 (Figure 2B), thus overcoming the limitations of polyploid cell lines where genome editing often results in a combination of correctly and incorrectly edited alleles (Canaj et al., 2019).

To obtain clonal cell lines with the desired modification, single SGFP2-positive cells were isolated by FACS sorting (Figure 2C-E). Because HAP1 cells have the tendency to become diploid or polyploid over time (Essletzbichler et al., 2014; Yaguchi et al., 2018), we used a gating strategy to specifically select for haploid cells (Figure 2 supplement 1). Genome editing of wild-type HAP1 (HAP1<sup>WT</sup>) cells with a targeting gRNA and the SGFP2-CTNNB1 repair construct resulted in a slight increase in fluorescence for a small population (0.2%) of cells compared to non-repaired control cells (Figure 2C-D). Treatment of these cells with CHIR99021, a potent and selective GSK3 inhibitor (Bain et al., 2007), resulted in an increase of the green fluorescence intensity (Figure 2E), but not the absolute number of positive events. The responsiveness to CHIR99021 provided a strong indication that these fluorescent
events corresponded to HAP1 cells in which the SGFP2 sequence was successfully knocked into the endogenous CTNNB1 locus (HAP1SGFP2-CTNNB1).

PCR based screening confirmed that 22/23 single-cell sorted clones indeed showed an integration of the expected size at the CTNNB1 locus. Complete sequence coverage of the insertion site in exon 2 was obtained for 9/11 sequenced clones, of which 8 showed the desired repair, and 1 clone showed an additional point mutation in the repaired locus. Sanger sequencing results around the SGFP2 integration site are shown for three correctly targeted HAP1SGFP2-CTNNB1 clones from three independent transfections, which were used for further experiments (Figure 2 supplement 2B-C). Thus, while scarless tagging of endogenous genes in HAP1 cells is relatively cumbersome (only 0.2% gated events), the desired repair occurs with almost 90% efficiency within this population.
Figure 2: Generation of HAP1^{SGFP2-CTNNB1} cell lines. A) Cartoon depicting exon 2 of the CTNNB1 locus, which contains the start codon, and the CTNNB1 protein before (left) and after (right) introduction of the SGFP2 by CRISPR/Cas9 mediated homology directed repair. B) Schematic of the experimental workflow and timeline for generating HAP1^{SGFP2-CTNNB1} clones. Cas9, gRNA and repair templates are transfected as plasmids. The repair template contains the coding sequence of SGFP2 surrounded by 800 bp homology arms on either side and lacks the gRNA recognition site (see supplement 2 of this figure). A short puromycin selection step is included from 24-48 hours after transfection to enrich for transfected cells. Haploid, GFP-positive cells are sorted and single cell clones are expanded for further analysis. C-E) FACS plots illustrating control (C) and SGFP2-CTNNB1 tagged cells (D-E). C) Cells transfected with Cas9 and gRNA in the absence of a repair template were used to set the gate for SGFP2-positive events. D) A small population of cells expressing low levels of SGFP2 can be detected when cells are transfected with Cas9, gRNA and repair template. E) Treatment of cells similar to those depicted in (D) with 8μM CHIR99021 does not change the amount of cells that are SGFP2 positive, but increases the SFP2 signal, most likely reflecting an increase in SGFP2-tagged beta catenin levels on a per cell basis and supporting the notion that the gated events indeed represent successfully tagged cells.
Figure 2 – supplement 1: FACS Gating strategy for haploid HAP1 cells. A-C) Single-cell gating based on forward scatter (FSC) and side scatter (SSC). D) Live cell gating based on DAPI exclusion. E-F) Haploid cell sorting based on Vibrant live-cell DNA dye. E) Haploid cell cycle profile. Only cells in G1 can be confidently identified as haploid (1n). The second peak contains both G2/M haploid cells, as well as diploid (2n) and polyplod events. Of note, the depicted HAP1WT population is mainly haploid. F) Back-gating of the haploid G1 population from E onto the forward and side scatter plot. A stringent gate is set based on cell size to ensure only G1 (1n) cells qualify for sorting.
Figure 2 – Supplement 2: SGFP2-CTNNB1 locus A) Detailed view of CTNNB1 exon 2 depicting gRNA design relative to the wildtype (top) and repaired (bottom) CTNNB1 allele. Note that the repair template contains the same sequence as the repaired allele depicted here. CTNNB1 sequences are shown in capital letters, SGFP2 sequences shown in lowercase. 5' UTR, SGFP2 and CTNNB1 and intron regions are indicated below the colored boxes. The gRNA (white arrow box above sequence) overlaps the start codon (depicted in bold), resulting in a Cas9-mediated double-strand break in the 5' UTR (predicted cut site indicated by dotted line and scissor, PAM site underlined). After successful homologous recombination, most of the gRNA binding site is destroyed, thus minimizing the chance of cutting the repair template or cutting the repaired allele. B-C) Sequencing of three independent HAP1-SGFP2-CTNNB1 clones on the 5' (B) and the 3' (C) end of SGFP2 integration in exon 2 of CTNNB1. Sanger sequencing of the endogenous CTNNB1 locus of clone 1, clone 2 and clone 3 shows an exact match to the design and thus correct homology directed repair.
To verify that the SGFP2 tag did not interfere with CTNNB1 function, three clonal HAP1<sup>SGFP2-CTNNB1</sup> cell lines were further characterized using established experimental readouts for WNT/CTNNB1 signaling (Figure 3 and Figure 3 supplement). Western blot analysis confirmed that the HAP1<sup>SGFP2-CTNNB1</sup> clones did not contain any untagged CTNNB1 but only expressed the fusion protein, which ran at the expected height (~27 kDa above the wild-type CTNNB1, Figure 3A). Moreover, in response to CHIR99021 treatment the total levels of SGFP2-CTNNB1 in tagged cell lines increased to the same extent as wild-type CTNNB1 in untagged cells, as detected with both CTNNB1 and GFP antibodies (Figure 3A-B). To examine if SGFP2-CTNNB1 was able to promote TCF/LEF transcriptional activity, we measured the induction of the TCF/LEF-responsive MegaTopflash luciferase reporter (Hu et al., 2007). Stimulation of the WNT/CTNNB1 pathway with CHIR99021 resulted in a similar range of reporter gene activation in the three tagged clones and parental HAP1 cells (Figure 3C). Induction of the universal WNT/CTNNB1 target gene AXIN2 (Lustig et al., 2002) was also comparable across wildtype and tagged cells (Figure 3D). Finally, we visualized the subcellular accumulation of SGFP2-CTNNB1 accumulation after WNT/CTNNB1 pathway activation. While untreated cells mainly show membrane localization of CTNNB1, treatment with purified WNT3A protein (Figure 3E) and CHIR99021 (Figure 3 supplement 1E) increased SGFP2-CTNNB1 levels in the cytoplasmic and nuclear.

Taken together, WNT-responsive changes in CTNNB1 levels and localization and activity are preserved after CRISPR/Cas9 mediated homozygous tagging of CTNNB1. Although there is some variation between the three clones with respect to CTNNB1 stabilization and target gene activation, this is likely due to the sub-cloning of these cell lines rather than the targeting
Based on the combined characterization results, we selected clone 2 for more extensive functional imaging experiments and analyses.

**Figure 3:** Functional validation of three independent HAP1<sup>SGFP2-CTNNB1</sup> clones. A) Western blot, showing CTNNB1 (HAP1<sup>WT</sup>) and SGFP2-CTNNB1 (HAP1<sup>SGFP2-CTNNB1</sup> clone 1, 2 and 3) accumulation in response to CHIR99021 treatment. All panels are from one blot that was cut at the 70 kDa mark and was stained with secondary antibodies with different fluorophores for detection. Top: HAP1<sup>WT</sup> cells express CTNNB1 at the expected wild-type size. Each of the three clonal HAP1<sup>SGFP2-CTNNB1</sup> cell lines only express the larger, SGFP2-tagged form of CTNNB1. Middle: Only the tagged clones express the SGFP2-CTNNB1 fusion protein, as detected with an anti-GFP antibody at the same height. Bottom: alpha-Tubulin (TUBA) loading control. A representative image of n=3 independent experiments is shown. B) Quantification of Western blots from n=3 independent experiments, including the one in (A), confirming that the accumulation of CTNNB1 in response to WNT/CTNNB1 pathway
activation is comparable between HAP1WT and HAP1SGFP2-CTNNB1 cells. Horizontal bar indicates the mean. C) Graph depicting the results from a MegaTopflash dual luciferase reporter assay, showing comparable levels of TCF/LEF reporter gene activation for HAP1WT and HAP1SGFP2-CTNNB1 cells in response to CHIR99021 treatment. Data points from n=3 independent experiments are shown. Horizontal bar indicates the mean. Values are depicted relative to the DMSO control, which was set to 1 for each individual cell line. D) Graph depicting AXIN2 mRNA induction in response to CHIR99021 treatment, demonstrating that induced expression of an endogenous target gene is comparable between HAP1WT and HAP1SGFP2-CTNNB1 cells. Data points represent n=3 independent experiments. Horizontal bar represents the mean. HPRT was used as a reference gene. Values are depicted relative to the HAP1WT DMSO control, which was set to 1.

E) Representative confocal microscopy images of the three HAP1SGFP2-CTNNB1 clones after 4-hour vehicle control or 100ng/ml WNT3A treatment, revealing intracellular accumulation of SGFP2-CTNNB1 (green). Nuclei were counterstained with SiR-DNA dye (magenta). Scale bar is 10μm.

Figure 3 supplement 1: Verification of the WNT/CTNNB1 responsiveness of HAP1 cells. A) Graph depicting AXIN2 qRT-PCR results from HAP1WT cells treated with the indicated range of CHIR99021 (1-10 μM) or DMSO vehicle control (0 μM) for 24 hours. HPRT was used as a reference gene. Error bars represent standard deviation within technical triplicates from n=1 biological experiment. Based on this, we selected 4μM and 8 μM as intermediate and high levels of WNT/CTNNB1 pathway induction for follow up experiments. B) Western blot, showing the increase in total (top) and non-phosphorylated (i.e. active) CTNNB1 levels (middle) in response to pathway stimulation. HAP1WT cells were treated for 24 hours with 4 or 8 μM CHIR99021, or DMSO vehicle control (0 μM). Alpha-Tubulin (TUBA, bottom) serves as a loading control. C-D) Quantification of the western blot from (B) depicting the relative fold change of total CTNNB1 (C) or non-phosphorylated CTNNB1 (D) to DMSO control corrected for Tubulin loading. E) Representative confocal microscopy images of three independent HAP1SGFP2-CTNNB1 clones, treated for 24 hours with 4 or 8 μM CHIR99021, or DMSO vehicle control. Scalebar is 10 μm.
Live imaging of endogenous SGFP2-CTNNB1 during WNT pathway activation

To better understand the temporal dynamics of endogenous CTNNB1 stabilization, we performed live-cell imaging over 12 hours in HAP1SGFP2-CTNNB1 cells (Figure 4, Supplementary Movie 1-3) with different levels of WNT stimulation. Unstimulated cells showed stable CTNNB1 localization at the cell membrane throughout the imaging time course (Figure 4A, Supplementary Movie 1). The membrane localization of CTNNB1 is consistent with its structural role in adherens junctions (Valenta, Hausmann, & Basler, 2012; Yap, Briehrer, & Gumbiner, 1997), which we will not consider further in the current study. Both WNT3A and CHIR99021 treatment resulted in a heterogeneous response pattern, with some cells in the population showing a far more prominent increase in CTNNB1 levels in the cytoplasm and nucleus than others (Figure 4A, Supplementary Movie 2-3).

To quantify these dynamic changes, we developed a custom-built automated segmentation pipeline in CellProfiler™ (Figure 4B). Quantification showed that the dynamics of CTNNB1 accumulation were independent of the dose of WNT3A (Figure 4 C-D, Supplementary Movies 4-5). Treatment with 100 ng/ml WNT3A increased SGFP2-CTNNB1 fluorescence 1.74-fold (mean, 95% CI 1.73-1.76) in the cytoplasm and 3.00-fold (mean, 95% CI 2.97-3.03) in the nucleus, with similar results in the other two HAP1SGFP2-CTNNB1 clones (Figure 4 supplement 1).

Treatment with 4 μM of CHIR99021 yielded similar kinetics as WNT3A treatment (Figure 4C-D). Treatment with WNT3A and 4 μM CHIR both also resulted in a decrease of SGPF2-CTNNB1 levels toward the end of the time series. However, at the highest levels of GSK3 inhibition (8 μM CHIR99021), no plateau was reached and the levels of SGFP2-CTNNB1 continued to increase throughout the imaging time course. The fact that intracellular SGFP2-CTNNB1 levels in the 8 μM CHIR99021 condition continued to accumulate, suggests that negative feedback,
for example through AXIN2 (Lustig et al., 2002), is overridden under these circumstances. Of note, the quantification also confirms that there is cell to cell heterogeneity in the response, regardless of whether WNT/CTNNB1 signaling is activated at the level of the receptor (WNT3A treatment) or at the level of the destruction complex (CHIR99021 treatment), as can be seen from the spread of intensities measured from individual cells (Figure 4 Supplement 2A-B).

Our quantification shows that, nuclear accumulation of CTNNB1 is favored over cytoplasmic increase (compare the slopes in Figure 4C-D). Moreover, the first significant increases in fluorescence intensity in the cytoplasm could be detected after ~45 minutes of treatment (Supplementary Movie 4, Figure 4 supplement 2C), whereas in the nucleus an increase was first after ~30 minutes (Supplementary Movie 5, Figure 4 supplement 2D). To examine the relation between the cytoplasmic and nuclear CTNNB1 pools more closely, we calculated the ratio between nuclear and cytoplasmic intensities of SGFP2-CTNNB1 (Figure 4E, Supplementary Movie 6). In untreated cells the nuclear/cytoplasmic ratio was 0.652 (mean, 95% CI 0.649-0.657), showing that SGFP2-CTNNB1 was preferentially localized to the cytoplasm (Figure 4E). For the first 3 hours after WNT3A and CHIR99021 addition, nuclear CTNNB1 levels rose considerably faster than cytoplasmic CTNNB1 levels until the nuclear/cytoplasmic ratio showed a slight nuclear enrichment of 1.08 (mean [3-5 hours] 95% CI 1.07-1.10) for 100 ng/ml WNT3A and 1.29 (mean [3-5 hours], 95% CI 1.26-1.32) for 8µM CHIR99021. This indicates that not only the turn-over, but also the subcellular localization of CTNNB1 is actively regulated both before and after WNT pathway activation.
Figure 4: Live imaging of HAP1-SGFP2-CTNNB1. A) Representative stills from confocal time-lapse experiments corresponding to Supplementary Movies 1-3, showing an increase of SGFP2-CTNNB1 after treatment with 100 ng/ml WNT3A (middle) and 8µM CHIR99021 (bottom) relative to a vehicle control (BSA) treated sample (top). Scale bar = 20 μm. B) Segmentation of nuclei (top) and cytoplasm (bottom) based in the SiR-DNA signal and SGFP2-CTNNB1 signal. Scale bar = 20μm. C-E) Quantification of time-lapse microscopy series, using the segmentation pipeline shown in (B). Arrow indicates the moment of starting the different treatments (T, see legend in E for details). C-D) Graph depicting the normalized intensity of SGFP2-CTNNB1 in the cytoplasm (C) or nucleus (D) over time. Solid lines represent the mean normalized fluorescence intensity and shading indicates the 95% confidence interval. n=155-400 cells for each condition and time point, pooled data from n=3 independent biological experiments. E) Graph depicting the nuclear/cytoplasmic ratio of SGFP2-CTNNB1 over time, calculated from raw intensity values underlying (C) and (D).

Figure 4 supplement 1: Graphs showing quantification of time-lapse microscopy experiments with three independent HAP1-SGFP2-CTNNB1 clones. Stills of this experiment are shown in Figure 3C. Segmentation was performed as described in Figure 4. Arrow indicates the moment of starting the different treatments (BSA in red or 100 ng/ml WNT3A in blue). Solid lines represent the mean normalized intensity and shading the 95% confidence interval in the cytoplasm (A) or nucleus (B). Line pattern indicates the three different clones. n=13-158 cells for each condition and time point for n=1 biological experiment.
Figure 4 supplement 2 A-B) Plots depicting the relative intensity (left) and the difference in relative intensity to BSA treated cells (right) in the cytoplasm (A) and nucleus (B) after 4 hours of treatment. Circles indicate the median value and bars indicate the 95% CI. In the relative intensity plot (left) the distribution is built from individual data points in a violin-type fashion to faithfully represent the distribution of data. In the difference plot (right) the distribution of differences is represented in a half violin plot. If the 95% CI in the difference plot does not overlap the zero line, which indicates no difference, the sample is significantly different from BSA control condition. C-D) Plots depicting the difference in relative intensity in the cytoplasm (C) and nucleus (D) between the moment of addition and 1 hour of treatment. Titles indicate the time (hh:mm). The distribution of differences is represented in a half violin plot. Circles indicate the median value and bars indicate the 95% CI. If the 95% CI does not overlap the zero line, which indicates no difference, the sample is significantly different from the BSA control condition.
Establishing a fitting model for SGFP2-CTNNB1 diffusion

Having measured the relative changes in the cytoplasmic and nuclear levels of CTNNB1 in response to WNT pathway activation, we next sought to exploit our experimental system to quantify additional molecular properties of CTNNB1 in each of these subcellular compartments using Fluorescence Correlation Spectroscopy (FCS). FCS is a powerful method to measure the mobility and absolute levels of fluorescent particles in a nanomolar range, compatible with typical levels of signaling proteins in a cell (reviewed in Hink, 2014). It has for instance been used to gain insight into the assembly of DVL3 supramolecular complexes (Yokoyama, Markova, Wang, & Malbon, 2012), the endogenous concentrations and mobility of nuclear complexes (Holzmann et al., 2019; Lam et al., 2012), and most recently, to quantify ligand-receptor binding reactions in the WNT pathway (Eckert et al., 2020). In point FCS, the fluorescence intensity is measured in a single point (Figure 5 A, D-E). Diffusion of a labeled particle, in this case SGFP2-CTNNB1, causes fluctuation of the fluorescence signal over time (Figure 5B). By correlating the fluorescence intensity signal to itself over increasing time-intervals, an autocorrelation curve is generated (Figure 5C). To extract relevant biophysical parameters, such as mobility (a measure for size) and the absolute numbers of the fluorescent particles, this autocorrelation curve is fitted with an appropriate model.

We first attempted to fit the autocorrelation curves obtained with point FCS measurements of HAP1SGFP2-CTNNB1 cells, with a model containing one single diffusion speed for SGFP2-CTNNB1. This model was unable to fit most of our data (Figure 5F). The current literature suggests that while a large portion of CTNNB1 is present as a monomer (Gottardi & Gumbiner, 2004; Maher, Mo, Flozak, Peled, & Gottardi, 2010), CTNNB1 is also present in multiprotein complexes in the cytoplasm and in the nucleus (reviewed in Gammons and Bienz, 2018).
Therefore, we next used a model with two diffusion components, in which the first diffusion component was fixed to the diffusion speed of monomeric, unbound SGFP2-CTNNB1 (14.9 µm²/s) and the second diffusion component was limited to slower speeds compatible with point-FCS imaging (see materials and methods for details). This model fitted well with our autocorrelation curves as obtained in both cytoplasmic and nuclear point FCS measurements (Figure 5G), consistent with the presence of free monomeric CTNNB1 and larger CTNNB1 containing complexes in both the nucleus and cytoplasm.
Figure 5: Two diffusion-component fit-model for SGFP2-CTNNB1 FCS measurements. A) Schematic representation of the point FCS technique, depicting the confocal volume with fluorescent particles diffusing in and out. Particles in FCS are defined by their coherent movement; therefore, a particle can be made up of monomers or multimers in isolation or complexed to unlabeled molecules. B) Schematic representation of intensity fluctuations over time as measured in the confocal volume. Fluctuations are the result of both photo-physics (e.g. blinking of the fluorophore) and diffusion. C) Graphical representation of the two diffusion-component fitting model used for our autocorrelation curves. $T_{\text{trip}}$ describes the blinking of the SGFP2 fluorophore and the after-pulsing artefact. $T_{\text{diff1}}$ and $T_{\text{diff2}}$ describe the monomeric and complexed form of SGFP2-CTNNB1, respectively. Details of all fitting parameters are described in Materials and Methods. D) Representative confocal images of HAP1 $^{15}$SGFP2-CTNNB1 cells treated for 4 hours with BSA (left) or 100 ng/ml WNT3A (right). E) Zoom in of the white rectangle in (D), with representative locations of FCS measurement points for cytoplasm (C) and nucleus (N) indicated with white crosses in the SGFP2-CTNNB1 channel and transmission channel. F-G) Fitting of a representative autocorrelation curve with one unfixed diffusion-component (F) or a two diffusion-component model (G), where the first diffusion component was fixed to the speed of free monomeric SGFP2-CTNNB1 (14.9 $\mu$m$^2$/s) and the second diffusion component was unfixed. The black line represents the autocorrelation curve generated from the FCS measurement, the red line represents the fitted model. The residuals after fitting of 25 individual curves are shown in the upper right corner of the graphs.
Quantification of SGFP2-CTNNB1 mobility in the nucleus

By fitting our data with this two-component diffusion model, we obtained the diffusion coefficient for the slower (i.e. complexed) CTNNB1 fraction in both the nuclear and cytoplasmic compartment. In the nucleus this second diffusion coefficient was 0.140 μm²s⁻¹ (median, 95%CI 0.121-0.213) in unstimulated cells and 0.178 μm²s⁻¹, (median, 95%CI 0.139-0.211) in cells treated with purified WNT3A (Figure 6A). This is comparable to the diffusion coefficients measured for other chromatin bound transcriptional activators (Lam et al., 2012), which could imply that this pool of SGFP2-CTNNB1 is linked to the chromatin as part of the TCF transcriptional complex (also called the “enhanceosome”). The fraction of SGFP2-CTNNB1 molecules bound to this nuclear complex increases from 0.26 (median, 95% CI 0.40-0.42) to 0.32 (median, 95%CI 0.30-0.35) upon WNT3A treatment (Figure 6B). This suggests that even in the absence of WNT3A a fraction of CTNNB1 might already be associated with the DNA and that WNT3A further stimulates the chromatin binding of CTNNB1, most likely through the TCF transcriptional complex.

Quantification of SGFP2-CTNNB1 mobility in the cytoplasm

In the cytoplasm, we determined the second diffusion coefficient of SGFP2-CTNNB1 to be 0.134 μm²s⁻¹ (median, 95% CI 0.124-0.196) in the absence of WNT3A stimulation (Figure 6A). This is indicative of very large complexes containing SGFP2-CTNNB1 that move with diffusion kinetics comparable to those previously observed for the 26S proteasome (C. G. Pack et al., 2014). After WNT3A treatment, the speed of the cytoplasmic complex increased 3.4-fold to 0.461 μm²s⁻¹ [95% CI of the median 0.367-0.573], indicating that the size of the cytoplasmic CTNNB1 complex changes when the WNT pathway is activated. However, the fraction of
CTNNB1 that is bound to a cytoplasmic complex remains largely unaltered upon WNT/CTNNB1 pathway stimulation (Figure 6B).

Determining the multimerization status of SGFP2-CTNNB1

Recent work suggests that the CTNNB1 destruction complex (also known as the “degradosome”) is a large and multivalent complex, mainly as the result of AXIN and APC multimerization (reviewed in Schaefer and Peifer, 2019). The second diffusion coefficient determined by our FCS measurements, is consistent with this model. Such a large, multivalent destruction complex would be expected to have multiple CTNNB1 binding sites. To measure the multimerization status (i.e. the number of bound SGFP2-CTNNB1 molecules) within this cytoplasmic complex, we performed Number and Brightness (N&B) analysis. N&B is a fluorescence fluctuation spectroscopy technique similar to point FCS, but it makes use of image stacks acquired over time rather than individual point measurements (Digman, Dalal, Horwitz, & Gratton, 2008). By quantifying the variance in fluorescence intensity of this stack, not only the number of particles but also their brightness can be determined. Because brightness is an inherent property of a fluorophore, a change in brightness is a measure of the number of fluorophores per particle. In our case, the brightness is indicative of the number of SGFP2-CTNNB1 molecules per complex. Unfortunately, N&B does not incorporate diffusion kinetics. Therefore, we cannot differentiate between monomeric (which would have a brightness of one) and complexed CTNNB1 (which would have a brightness exceeding one if multiple CTNNB1 molecules reside in a single complex). Therefore, the measured brightness of SGFP2-CTNNB1 in our N&B analysis is the average of both fractions. We observe that the total pool of SGFP2-CTNNB1 has a brightness similar to EGFP and SGFP2 monomers in both the cytoplasm and nucleus (Figure 6C). This suggests that few, if any, of the cytoplasmic or
nuclear complexes measured by point FCS, contain multiple SGFP2-CTNNB1 molecules. If the
cytoplasmic SGFP2-CTNNB1 containing complex indeed represents a large, multivalent
destruction complex, this would imply that, quite unexpectedly, most CTNNB1 binding sites
are unoccupied in both the absence and presence of WNT3A.

**Quantification of SGFP2-CTNNB1 concentrations**

FCS also allowed us to determine the absolute number of SGFP2-CTNNB1 molecules in the
confocal volume (Figure 6D). After calibration of the confocal volume via reference
measurements, the particle number can be converted to the concentration of endogenous
SGFP2-CTNNB1 in the cell (Table 1). In the absence of WNT, the concentration of SGFP2-
CTNNB1 was 202 nM (median, 95%CI 151-253) in the cytoplasm and 144 nM (median, 95%CI
111-177) in the nucleus. This is consistent with the nuclear exclusion we observed with
confocal imaging (Figure 4E). We measured comparable particle numbers with N&B (Figure 6
supplement 1A).

In the presence of WNT3A, we measure a 1.1-fold increase in the total SGFP2-CTNNB1
concentration to 219 nM (median, 95%CI 173-264 nM) in the cytoplasm. This increase is
smaller than expected from fluorescence intensity measurements (Figure 4C, Figure 6
supplement 1), for which we currently have no explanation. In the nucleus the concentration
increases 2.1-fold to 300 nM (median, 95%CI 257-342) upon pathway activation. Nuclear
concentrations of SGFP2-CTNNB1 therefore exceed cytoplasmic concentrations after WNT3A
treatment, consistent with the nuclear accumulation observed with live imaging (Figure 4).
Our observed concentrations of endogenous CTNNB1 in HAP1 cells are in a similar range as
those previously determined by quantitative mass spectrometry in different mammalian cell
lines (Kitazawa et al., 2017; Tan et al., 2012). Of note, the exact concentrations can vary
between cell types and may be dependent on the intricacies and assumptions of each individual technique.

Using the fractions obtained from the FCS fitting (Figure 6B), we also determined the concentration of fast (i.e. free monomeric) and slow (i.e. complexed) SGFP2-CTNNB1 (Figure 6E-F, Table 2). In the cytoplasm we see that the concentration of both fast and slow SGFP2-CTNNB1 increases upon WNT3A treatment. This challenges the view that mainly monomeric CTNNB1 accumulates, as commonly depicted (Figure 1). In the nucleus, the concentration of fast moving CTNNB1 increases 2.0-fold from 95 nM (median, 95%CI 85-133) to 186 nM (median, 95%CI 164-238), while slow moving CTNNB1 concentrations increase 4.4-fold from 23 nM (median, 95%CI 4-43) to 102 nM (median, 95%CI 67-114). The preferential increase of the slow-moving fraction suggests that binding affinity of CTNNB1 to the chromatin, and presumably TCF transcriptional complexes, is increased after WNT3A treatment.

Taken together, this is the first time that the concentrations of different functional pools of endogenous CTNNB1 have been measured in living mammalian cells using functional imaging.
Figure 6: Mobility and abundance of SGFP2-CTNNB1 molecules in living cells after 4 hours WNT3A treatment or control. A-B) Graphs depicting the speed (A) and fraction (B) of the second diffusion component (i.e. SGFP2-CTNNB1 containing complex) measured by FCS. C) Graph depicting the molecular brightness of SGFP2-CTNNB1 in the cytoplasm and nucleus relative to controls as measured with N&B in the
same subcellular compartments. EGFP monomer was used for normalization and EGFP dimer as a control for N&B measurements. D) Graph depicting the total number of SGFP2-CTNNB1 particles (monomeric plus complexed) as measured with FCS. E) Graph depicting the number of SGFP2-CTNNB1 particles with the fast diffusion component (i.e. free monomeric). F) Graph depicting the number of SGFP2-CTNNB1 containing particles with the slow diffusion component (i.e. complex associated).

**Figure 6 supplement 1**

A) Graph depicting the total number of SGFP2-CTNNB1 particles as measured with N&B. B) Graph depicting the average fluorescence intensity at the start of the FCS measurement. The increase in SGFP2-CTNNB1 fluorescence in the cytoplasm (2.1-fold) exceeds the increase in the SGFP2-CTNNB1 concentration (1.1-fold, Figure 6D), but does correspond to the relative increase measured by time-lapse imaging (1.7-fold, Figure 4C). D) Graph depicting fluorescence lifetimes calculated from the FCS measurements. The Fluorescence lifetime of SGFP2-CTNNB1 is independent of the subcellular compartment and treatment. Therefore, photophysical effects are not the cause for the difference between the fold-change in fluorescence and concentrations of the FCS measurements as described in C).

**Table 1:** Total number of SGFP2-CTNNB1 molecules and calculated concentrations obtained with FCS

| compartment | treatment | Number of molecules | Concentration (nM) |
|------------|-----------|---------------------|--------------------|
|            |           | median 95% CI       | median 95% CI      |
| Cytoplasm  | BSA       | 80 67-113           | 202 151-253        |
|            | WNT3A     | 95 84-121           | 219 173-264        |
|            | BSA       | 63 53-71            | 144 111-177        |
|            | WNT3A     | 135 128-150         | 300 257-342        |
| Nucleus    | BSA       | 45 37-63            | 95 85-133          |
|            | WNT3A     | 97 81-100           | 186 164-238        |

**Table 2:** Number and concentration of SGFP2-CTNNB1 molecules with fast or slow diffusion times obtained with FCS

| compartment | treatment | Number of molecules | Concentration (nM) |
|------------|-----------|---------------------|--------------------|
|            |           | median 95% CI       | median 95% CI      |
| Cytoplasm  | BSA       | 49 41-65            | 104 70-149         |
|            | WNT3A     | 60 46-78            | 145 84-181         |
|            | BSA       | 45 37-63            | 95 85-133          |
|            | WNT3A     | 97 81-100           | 186 164-238        |
| Nucleus    | BSA       | 60 42-79            | 21-37              |
|            | WNT3A     | 75 63-91            | 29-41              |
|            | BSA       | 23 4-42             | 2-24               |
|            | WNT3A     | 102 67-114          | 39-51              |
A minimal computational model of WNT/CTNNB1 signaling

Quantitative measurements and physical parameters of WNT pathway components and their interactions remain limited (Kitazawa et al., 2017; Lee, Salic, Krüger, Heinrich, & Kirschner, 2003; Tan et al., 2012), especially in living cells. As we obtained measurements of different functional pools of CTNNB1, we next sought to integrate these parameters in a minimal computational model of WNT signaling to identify the critical nodes of regulation of subcellular SGFP2-CTNNB1 (Figure 7A, Table 3-4, Materials and Methods). This minimal model is based on a previous model by Kirschner and colleagues (Lee et al., 2003), and incorporates the new data obtained in our study, supplemented with parameters from literature (Lee et al., 2003; Tan et al., 2012).

Our model diverges from the model presented by Lee et al. on two major points. First, the model is simplified by omitting the details of the destruction complex cycle and the action of APC and AXIN, since our data do not provide new quantitative data on this part of the WNT/CTNNB1 signaling cascade. Second, we explicitly include shuttling of CTNNB1 between the cytoplasm and nucleus in both directions (Schmitz, Rateitschak, & Wolkenhauer, 2013; Tan, Gardiner, Hirokawa, Smith, & Burgess, 2014).

Thus, our model (Figure 7A) describes the binding of cytoplasmic CTNNB1 (‘CB’) to the destruction complex (‘DC’) leading to its phosphorylation and degradation, which releases the DC. Transport of CTNNB1 from the cytoplasm to the nucleus, allows nuclear CTNNB1 (‘NB’) to bind to TCF forming a transcriptional complex (NB-TCF). When WNT is present in the system, we describe the inactivation of the destruction complex (‘DC*’) by DVL.

Our final model faithfully recapitulates the dynamic changes that we observe with functional imaging (compare Figure 7B-F to Figure 4 and 6). Moreover, it reveals two critical regulatory
nodes in addition to the requisite inactivation of the destruction complex. The first additional
node of regulation is nuclear import and export (or ‘shuttling’, described by $k_6/k_7$). Upon WNT
stimulation, the ratio of $k_6/k_7$ needs to increase in order for the model to match the free
CTNNB1 concentrations we measured by FCS (Table 4, Figure 6E). Thus, the balance shifts
from nuclear export before WNT, to nuclear import after WNT. The second additional node
of regulation is the association of CTNNB1 with the TCF transcriptional complex (or
‘retention’), described by $k_9/k_8$. Upon WNT stimulation, the ratio of $k_9/k_8$ needs to decrease
by more than a factor of 10 in order for the model to reproduce the concentrations of free
and bound CTNNB1 in the nucleus as measured by FCS (Table 4, Figure 6E-F). Thus, association
of CTNNB1 to the TCF transcriptional complex is favored after WNT stimulation.

In summary, our model suggests that WNT/CTNNB1 signaling is regulated at three distinct
levels of the signal transduction pathway: destruction complex inactivation,
nucleocytoplasmic shuttling and nuclear retention.
Figure 7: Computational model of WNT/CTNNB1 based on FCS for free and complexed CTNNB1 (Table 1-2). A) Schematic overview of the model. DC=destruction complex, DC*=DVL-inactivated DC, CB=cytoplasmic CTNNB1, CB*=phosphorylated CB, NB=nuclear CTNNB1, TCF=TCF/LEF transcription factors, DVL=WNT-activated DVL. The model assumes that there is no activated DVL in the absence of WNT, therefore k₄/k₅ do not play any role in the WNT ‘OFF’ equilibrium. Note that CB* is degraded and therefore plays no role in the model. B) Graph depicting the modelled concentrations of cytoplasmic components over time. The black line indicates total concentration of cytoplasmic CTNNB1, corresponding to Figure 4C. C) Graph depicting the modelled concentrations of nuclear components over time. The black line indicates total concentration of nuclear CTNNB1, corresponding to Figure 4D. D) Graph depicting the ratio of total nuclear and cytoplasmic CTNNB1 over time, corresponding to the measurements in 4E. E) Graph depicting the DC-bound CTNNB1 fraction ratio over time. F) Graph depicting the TCF-bound CTNNB1 fraction ratio over time.
Table 3: Variables Minimal Model of WNT signaling.

| Model name | Variable | Compound | Values obtained from | WNT ON (nM) | WNT OFF (nM) |
|------------|----------|----------|----------------------|-------------|-------------|
| CB         | $x_1$    | Free cytoplasmic CTNNB1 | FCS data this report | 104         | 145         |
| DC         | $x_2$    | Free destruction complex | Model equations     | 77.3        | 55.5        |
| CB*-DC     | $x_3$    | DC-bound phosphorylated CTNNB1 | FCS data this report* | 67*         | 67*         |
| DC*        | $x_4$    | Inactivated destruction complex | Model equations     | 0           | 21.6        |
| NB         | $x_5$    | Free nuclear CTNNB1 | FCS data this report | 95          | 186         |
| TCF        | $x_6$    | Free TCF | Model equations      | 99          | 20          |
| NB-TCF     | $x_7$    | TCF-bound nuclear CTNNB1 | FCS data this report | 23          | 102         |
| TCF0       | --------- | Total TCF | $x_5$ and Tan et al., 2012 - Figure 11 | 122         | 122         |

*Under the assumption that $k_3$ does not change, the levels of CB*-DC remain equal. Since there was no significant difference between the concentration of slow SGFP2-CTNNB1 (Table 2) the average of both medians was used.

Table 4: Equilibrium conditions for the Minimal Model of WNT signaling. All rates are multiplied with factor R=20, so that the equilibrium is reached at 4.5h according to Figure 4 C, D.

| Rate constant | Biological process | Values based on | WNT OFF | WNT ON |
|---------------|--------------------|-----------------|---------|--------|
| $b$           | nM/min$^1$         | CTNNB1 synthesis | $v_{12}$ from Lee | 0.423  | 0.423  |
| $k_2$/$k_1$   | nM                 | Binding to and phosphorylation by the destruction complex of cytoplasmic CTNNB1 | $K_b$ from Lee | 120    | 120    |
| $k_3$         | min$^{-1}$         | Dissociation and degradation of phosphorylated CTNNB1 from the destruction complex | Deduced from $b$ and $x_3$ | 0.0063 | 0.0063 |
| $k_4$/$k_4^*$ | nM                 | Inactivation of the destruction complex by activated DVL | Fitted to $x_1$ and $x_7$ | N.A.   | 2.56   |
| $k_5$/$k_7^*$ |                    | Ratio between nuclear import and export of CTNNB1 | Deduced from $x_1$ and $x_5$ | 0.913  | 1.28   |
| $k_6$/$k_8$   | nM                 | Dissociation of nuclear CTNNB1 from TCF | Deduced from $x_6$, TCF0, $x_7$ | 409    | 36.5   |
Discussion

WNT signaling is a critical pathway for tissue development and homeostasis. Although most core players and many of their molecular interaction mechanisms have been uncovered, dynamic spatiotemporal information at the endogenous level and with subcellular resolution is very limited. As both genome editing approaches and quantitative live-cell microscopy have advanced further, the goal of studying WNT/CTNNB1 signaling at endogenous expression levels in living cells now is within reach. Maintaining endogenous expression levels is important, as overexpression may lead to altered stoichiometry of signaling components, as well as changes in subcellular localization (T. J. Gibson et al., 2013; Mahen et al., 2014). Indeed, it has been shown that exogenously expressed CTNNB1 is less signaling competent, probably due to its post-translational modification status (Hendriksen et al., 2008). Here we generated and characterized clonal HAP1*SGFP2-CTNNB1 knock-in cell lines to study the dynamic behavior and subcellular complex state of endogenous CTNNB1 in individual living human cells in the presence and the absence of WNT pathway activation (Figure 2-3).

Using live-cell microscopy and automated cell segmentation we observe that endogenous CTNNB1 only increases 1.7-fold in the cytoplasm and 3.0-fold in the nucleus after WNT3A treatment, which is consistent with the literature (Jacobsen et al., 2016; Kafri et al., 2016; Massey et al., 2019). Our data show a faster increase of endogenous CTNNB1 upon GSK3 inactivation or WNT stimulation than two recent studies that used chromobody labeling (Keller et al., 2018; Traenkle et al., 2015). This could be a cell type specific difference, but might also be related to the background expression levels of unbound chromobodies that could mask subtle and early changes in endogenous protein levels. In addition, we show that the response to WNT3A stimulation and even to global GSK3 inhibition is heterogeneous (Figure 4A, Figure 4 supplement 2A-B). There are indications that cell cycle, adhesion status...
and mechano-signaling are important in determining the WNT response (Benham-Pyle, Pruitt, & Nelson, 2015; Benham-Pyle, Sim, Hart, Pruitt, & Nelson, 2016; Gayrard, Bernaudin, Déjardin, Seiler, & Borghi, 2018; Howard, Deroo, Fujita, & Itasaki, 2011; Kafri et al., 2016; Olmeda, Castel, Vilaró, & Cano, 2003; van Leeuwen, Byrne, Jensen, & King, 2007). Although, it will be interesting to use live-cell imaging to further uncover the roles of these factors in determining the cell-to-cell variability in WNT responsiveness in the future, here we focus on novel regulatory aspects of CTNNB1 in the cytoplasm and nucleus.

Cytoplasmic regulation of CTNNB1

Using FCS, we show for the first time that CTNNB1 resides in slow moving complexes in the cytoplasm of living mammalian cells (Figure 5-6). The main known cytoplasmic complex containing CTNNB1 is the destruction complex. The combined weight of the individual destruction complex components (AXIN, APC, CSNK1 and GSK3) would be expected to result in a much higher mobility than the cytoplasmic CTNNB1-containing complex we observed. Currently, evidence is growing that the destruction complex forms phase separated aggregates (also termed biomolecular condensates) (reviewed in Schaefer and Peifer, 2019). Oligomerization of AXIN and APC underlies the formation of these aggregates, and this in turn appears to be required for efficient degradation of CTNNB1 (Fiedler, Mendoza-Topaz, Rutherford, Mieszczanek, & Bienz, 2011; Kunttas-Tatli, Roberts, & McCartney, 2014; Pronobis, Deuitch, Posham, Mimori-Kiyosue, & Peifer, 2017; Spink, Polakis, & Weis, 2000). There is some evidence that these aggregates form at (near) endogenous levels (Fagotto et al., 1999; Faux et al., 2008; Mendoza-Topaz, Mieszczanek, & Bienz, 2011; Pronobis, Rusan, & Peifer, 2015; Schaefer et al., 2018; Thorvaldsen et al., 2015), but it is still an open question what the exact composition and size of the destruction complex is in a physiological context. It should
be noted that our imaging does not visualize such aggregates (Figure 4A). Nevertheless, the diffusion coefficients we observed for the cytoplasmic CTNNB1 complex are compatible with a large complex. Interestingly, our N&B data indicate that few, if any, complexes exist that contain multiple SGFP2-CTNNB1 molecules, which would logically be expected for a large oligomerized destruction complex with multiple CTNNB1 binding sites.

Alternative explanations for the slow diffusion speed of the cytoplasmic CTNNB1 complex include a monovalent destruction complex in association with (parts of) the proteasome machinery (Li et al., 2012; Lui et al., 2011; Schaefer et al., 2020) or anchoring of either CTNNB1 itself or any of its interaction partners to the cytoskeleton (reviewed in Bryja et al., 2017; Fagotto, 2013). Therefore, additional data on the localization and mobility of other endogenously tagged WNT pathway components are required to determine the exact identity of the slow-diffusing, cytoplasmic CTNNB1 complex.

Of note, our data clearly show that a substantial fraction of CTNNB1 in the cytoplasm remains bound upon pathway stimulation (Figure 6B). Additionally, we show that this complex has an increased mobility compared to control treated cells (Figure 6A). Therefore, while the diffusion coefficient is still very low (indicating a very large complex), this implies it is a different complex than that observed in the absence of WNT stimulation. The current literature suggests that the destruction complex is sequestered to the FZD-LRP receptor complex upon WNT pathway stimulation. Several models exist for how the membrane sequestration inhibits CTNNB1 degradation, including LRP mediated GSK3 inhibition (Stamos, Chu, Enos, Shah, & Weis, 2014), sequestration of GSK3 in multi vesicular bodies (Taelman et al., 2010), (partial) dissociation of the destruction complex (X. Liu, Rubin, & Kimmel, 2005; Tran & Polakis, 2012), and saturation of CTNNB1 within an intact destruction complex (Li et
None of these models include a remaining cytoplasmic CTNNB1 complex in the presence of WNT ligands, but our data suggest that such a complex must exist.

### Nuclear regulation of CTNNB1

The key function of CTNNB1 downstream of WNT is to regulate transcription of TCF/LEF target genes (Doumpas et al., 2019; Schuijers, Mokry, Hatzis, Cuppen, & Clevers, 2014). Proteomic analyses have shown that the WNT-responsive transcriptional complex consists of CTNNB1, TCF/LEF, PYGO and BCL9 and several other large proteins (Fiedler et al., 2015; van Tienen et al., 2017). Using FCS we showed that CTNNB1 resides in a nuclear complex with a diffusion coefficient that is compatible with such a DNA-bound transcriptional complex (Figure 6A) (Lam et al., 2012). Upon pathway activation we see an increase in the fraction and absolute levels of this slow-diffusing nuclear CTNNB1 complex, consistent with increased CTNNB1 binding to its target sites (Figure 6B, F). The concentration increases to a 102 nM, which corresponds to something in the order of 20,000 bound CTNNB1 molecules in one nucleus, assuming a small nuclear volume of 0.36 pL (Tan et al., 2012). Published CHIPseq studies report many CTNNB1 DNA binding sites, ranging from several hundred to several thousand sites in mammalian cells (Cantù et al., 2018; Doumpas et al., 2019; Schuijers et al., 2014). It is therefore conceivable that at least some of the slow-diffusing CTNNB1 particles we measure represent CTNNB1 that is associated with DNA bound TCF transcriptional complexes.

Although CTNNB1 is known to associate with TCF/LEF factors in response to WNT/CTNNB1 signaling to drive transcription (Franz, Shlyueva, Brunner, Stark, & Basler, 2017; Schuijers et al., 2014), we also detect low levels of nuclear CTNNB1 complex in the absence of a WNT stimulus (Figure 6B,F). The diffusion coefficient of the nuclear CTNNB1 complex does not change upon the addition of WNT3A, suggesting that some CTNNB1 is already associated with
the DNA even in the absence of a WNT stimulus. Whether the nuclear CTNNB1 complex, either in the absence or presence of WNT, truly represents TCF/LEF associated CTNNB1 remains to be determined. At this point, we cannot exclude the contribution to TCF/LEF independent DNA binding (Armstrong, Fagotto, Prothmann, & Rupp, 2012; Essers et al., 2005; Kormish, Sinner, & Zorn, 2010), or anomalous subdiffusion in the nucleus, either due to physical obstruction, transient DNA-binding events protein or protein complex formation (Dross et al., 2009; Kaur et al., 2013; Wachsmuth, Waldeck, & Langowski, 2000).

Regulation of CTNNB1 nuclear accumulation

In HAP1 cells, endogenous CTNNB1 is excluded from the nucleus in the absence of WNT. Our live imaging data reveal an immediate and preferential increase in nuclear CTNNB1 upon WNT3A stimulation, until an equilibrium is reached between the cytoplasmic and nuclear levels (Figure 4E, Figure 6D). This has previously been observed in HEK293 cells stably overexpressing low levels of YFP-CTNNB1 (Kafri et al., 2016). Indeed, our computational model can explain this preferential increase in nuclear CTNNB1 by a shift in the balance of nuclear translocation from nuclear export to nuclear import after the addition of WNT3A (Figure 7). This suggests that WNT signaling not only regulates the absolute levels of CTNNB1 through destruction complex inactivation, but also actively changes its subcellular distribution.

Intriguingly, CTNNB1 does not contain nuclear import or export signals and can translocate independently of classical importin and exporter pathways (Fagotto, Glück, & Gumbiner, 1998; Wiechens & Fagotto, 2001; Yokoya, Imamoto, Tachibana, & Yoneda, 1999). Hence, the molecular mechanism of CTNNB1 subcellular distribution remains incompletely understood.

Evidence from Fluorescence Recovery After Photobleaching (FRAP) studies suggest that the
increase in nuclear CTNNB1 is due to changes in binding to its interaction partners in the cytoplasm and nucleus (retention) rather than active changes in nuclear import and export rates (shuttling) (Jamieson, Sharma, & Henderson, 2011; Krieghoff, Behrens, & Mayr, 2006). We argue that the two are not mutually exclusive, as our experimental data and computational model suggest that WNT regulates both nucleocytoplasmic shuttling and nuclear retention of CTNNB1. Indeed, we see an increase of nuclear CTNNB1 complexes in the nucleus (Figure 6B, F) and the dissociation of CTNNB1 from TCF is reduced over 10-fold in WNT signaling conditions in our computational model (Table 4). This could be achieved through posttranslational modification and conformational changes in CTNNB1 as proposed by others (Gottardi & Gumbiner, 2004; Sayat, Leber, Grubac, Wiltshire, & Persad, 2008; Wu et al., 2008).

Challenges and opportunities for fluorescence fluctuation spectroscopy techniques

Using fluorescence fluctuation spectroscopy techniques (FCS and N&B) we have quantified endogenous CTNNB1 concentrations and complexes in living cells for the first time. This has yielded novel insights into the CTNNB1 complex state and new parameters for computational modelling. As with any technique, there are several limitations of point-FCS that we should consider.

First, we are limited by the assumptions we make in the FCS fitting model used. Although obvious mistakes in assumptions immediately become clear due to bad fitting results and can therefore be excluded, not every wrong assumption will stand out accordingly. Our data clearly shows that assuming only one diffusion speed for CTNNB1 in cells would be incorrect (Figure 5). However, whether we measure a single distinct large complex with the second diffusion speed, or rather an average of multiple different CTNNB1 containing complexes...
cannot be determined in our current set-up. In addition, we assume that CTNNB1 is present as a free-floating monomer (as fixed for our first component), based on previous observations (Gottardi & Gumbiner, 2004; Maher et al., 2010). However, at least one report suggests that CTNNB1 is not present as a monomer but rather in small cytoplasmic complexes of ~200 kDa (Gerlach, Emmink, Nojima, Kranenburg, & Maurice, 2014). As diffusion speed is relatively insensitive to differences in size (e.g. an 8-fold increase in protein mass is expected to result in only a 2-fold reduction of the diffusion coefficient for a spherical particle), it is possible that we do not measure truly free-floating CTNNB1, but rather smaller complexes.

Secondly, point FCS gives an indication of the size of the cytoplasmic and nuclear complexes we observe. However, it cannot provide conclusive evidence on their composition. An exciting possibility would be to label additional components presumed to be present in the CTNNB1-containing complexes at the endogenous level to uncover the precise composition and stoichiometry of protein complexes involved in WNT signaling. For instance Fluorescence Cross Correlation Spectroscopy (FCCS) could be employed to test if two proteins reside within the same complex (Elson, 2011; Hink, 2014; Macháň & Wohland, 2014). A combination of such functional imaging techniques, biochemical and proteomic approaches, together with specific perturbations and mutant versions of candidate proteins can be employed to further our understanding of the dynamic composition of endogenous CTNNB1 complexes, as well as to help us resolve how WNT signaling alters nucleocytoplasmic shuttling of CTNNB1 and specifically induces nuclear retention of CTNNB1.

Summary and outlook

In conclusion, we generated a functional fluorescent endogenous CTNNB1 fusion using CRISPR/Cas9-mediated genome editing. Using live imaging we show a preferential increase in
nuclear accumulation of CTNNB1 upon WNT pathway stimulation. Furthermore, state-of-the-art, quantitative microscopy revealed changes in the abundance and identity of nuclear and cytoplasmic CTNNB1 complexes upon WNT pathway activation. Combining our experimental data with computational modeling of WNT/CTNNB1 signaling reveals that WNT regulates the dynamic distribution of CTNNB1 across different functional pools not only by modulating the destruction complex, but also by actively changing nucleocytoplasmic shuttling and nuclear retention.

Quantitative functional analyses of proteins at endogenous levels will help us to further understand the mechanisms of WNT/CTNNB1 signaling and other cellular signal transduction pathways. As both genome editing and live cell imaging techniques continue to improve, additional possibilities will open up to address longstanding questions in cellular signaling in a physiological context with high spatial and temporal resolution. New opportunities and challenges await as these investigations extend to 3D organoid cultures, developing embryos and living organisms.
Material and Methods

DNA Constructs

The following constructs were used: pSpCas9(BB)-2A-Puro (PX459) V2.0 ((Ran et al., 2013), a kind gift from Feng Zhang, available from Addgene, plasmid #62988), MegaTopflash ((Hu et al., 2007), a kind gift from Dr. Christophe Fuerer and Dr. Roel Nusse, Stanford University), CMV Renilla (E2261, Promega, Madison, WI), pSGFP2-C1 ((G.-J. Kremers et al., 2007), a kind gift from Dorus Gadella, available from Addgene, plasmid #22881), pMScarlet-i_C1 (Bindels et al., 2017), a kind gift from Dorus Gadella, available from Addgene, plasmid #85044), pSYFP2-C1 ((G. J. Kremers et al., 2006), a kind gift from Dorus Gadella, available from Addgene, plasmid #22878), mTurquoise2-C1 ((Goedhart et al., 2012), a kind gift from Dorus Gadella, available from Addgene, plasmid #54842), pEGFP (Clontech, Mountain View, CA), pEGFP2 ((C. Pack, Saito, Tamura, & Kinjo, 2006), a kind gift from Masataka Kinjo) and pBluescript II KS(+) (Stratagene, La Jolla, CA).

The gRNA targeting the start codon in exon2 of human CTNNB1 was designed using the MIT webtool (crispr.mit.edu) and cloned into pX459. Oligos RVA567 and RVA568 (Table 5) encoding the gRNA were annealed, and ligated into BbsI-digested pX459 plasmid as previously described (Ran et al., 2013) to obtain pX459-CTNNB1-ATG.

The repair plasmid for SGFP2-CTNNB1 (pRepair-SGFP2-CTNNB1) was cloned using Gibson cloning (D. G. Gibson et al., 2009). First, a repair plasmid including the Kozak sequence from the pSGFP2-C1 plasmid was generated (pRepair-Kozak-SGFP2-CTNNB1). For this, 5’ and 3’ homology arms were PCR amplified from genomic HEK293A DNA with primers RVA618 and RVA581 (5’ arm) or RVA619 and RVA584 (3’ arm). SGFP2 was amplified with Gibson cloning from pSGFP2-C1 with primers RVA582 and RVA583 and the backbone was amplified from SacI.
digested pBlueScript KS(+) with primers RVA622 and RVA623. The final repair construct (pRepair-SGFP2-CTNNB1) contains the endogenous CTNNB1 Kozak sequence before the SGFP2 ATG. To obtain (pRepair-SGFP2-CTNNB1), the backbone and homology regions were amplified from pRepair-SGFP2-Kozak-CTNNB1 with primers RVA1616 and RVA1619 and an SGFP2 without the Kozak sequence was amplified from pSGFP2-C1 with primers RVA1617 and RVA1618. To generate color variants of the repair plasmid SYFP2, mScarlet-i and mTurquoise2 were also amplified from their respective C1 vectors with primers RVA 1617 and RVA 1618.

PCR products were purified and assembled with a Gibson assembly master mix with a 1:3 (vector:insert) molar ratio. Gibson assembly master mix was either purchased (E2611S, NEB) or homemade (final concentrations: 1x ISO buffer (100mM Tris-HCL pH 7.5, 10mM MgCl2, 0.2M dNTPs (R0181, Thermo Scientific), 10mM DTT (10792782, Fisher), 5% PEG-8000 (1546605, Sigma-Aldrich, St Louis, MO), 1mM NAD+ (B9007S, NEB)), 0.004 U/µl T5 exonuclease (M0363S, NEB), 0.5 U/µl Phusion DNA Polymerase (F-530L, Thermo Scientific) and 4 U/µl Taq DNA ligase (M0208S, NEB)).

The following plasmids are available from Addgene: pX459-CTNNB1-ATG (#153429), pRepair-SGFP2-CTNNB1 (#153430), pRepair-mScI-CTNNB1 (#153431), pRepair-SYFP2-CTNNB1 (#153432), pRepair-mTq2-CTNNB1 (#153433)).

Primers used

| Primer name | Sequence |
|-------------|----------|
| RVA24       | CAAGTTTGGTGTAGGATATGCC |
| RVA25       | CGATGTCAATAGGACTCCAGA |
| RVA124      | AGTGTGAGGTCCACGGAAA |
| RVA125      | CCGTCATGGACATGGGAAT |
| RVA567      | CACCGTGAGTAGCCATTGTCCACGC |
| RVA568      | AAACCGGTGGACAAATGGCTACTCAC |
| RVA581      | tgctcaccatgtggtATTTTCAAAACAGTTGTATGGTATACTTC |
| RVA582      | actgttttgaagaatcCCACCATGGTGAGCAAGGC |

| Table 5: primers/oligonucleotides used in this study |
Cell Culture, Treatment and Transfection

HAP1 cells (a kind gift from Thijn Brummelkamp, NKI) were maintained in full medium (colorless IMDM (21056023, Gibco, Thermo Fisher Scientific, Waltham, MA) supplemented with 10% FBS (10270106, Gibco) and 1X Glutamax (35050061, Gibco)) under 5% CO2 at 37°C in humidifying conditions and passaged every 2-3 days using 0.25% Trypsin-EDTA (25200056, Gibco). Cells were routinely tested for mycoplasma. We verified the haploid identity of the parental HAP1WT by karyotyping of metaphase spreads. To maintain a haploid population, cells were resorted frequently (see below) and experiments were performed with low passage number cells.

Where indicated, cells were treated with CHIR99021 (6mM stock solution in DMSO) (1677-5, Biovision, Milpitas, CA) or Recombinant Mouse Wnt-3a (10µg/ml stock solution in 0.1% BSA in PBS) (1324-WN-002, R&D systems, Bio-Technne, Minneapolis, MN) with DMSO and 0.1% BSA in PBS as vehicle controls, respectively.

Cells were transfected using Turbofect (R0531, ThermoFisher, Thermo Fisher Scientific, Waltham, MA), X-tremeGene HP (6366546001, Roche, Basel, Switzerland) or Lipofectamine
3000 (Invitrogen, Thermo Fisher Scientific, Waltham, MA) in Opti-MEM (Gibco) according to the manufacturer's instructions.

HAP1<sup>SGFP2-CTNNB1</sup> generation

800,000 HAP1 cells/well were plated on 6-well plates. The following day, cells were transfected with Turbofect and 2000 ng DNA. pX459-CTNNB1-ATG and pRepair-SGFP2-CTNNB1 were transfected in a 2:1, 1:1 or 1:2 ratio. pSGFP2-C1, pX459 or pX459-CTNNB1-ATG were used as controls. From 24 to 48 hours after transfection cells were selected with 0.75 µg/ml puromycin (A1113803, Gibco). Next, cells were expanded and passaged as needed until FACS sorting at day 9. For FACS analysis and sorting cells were washed, trypsinized, resuspended with full medium and spun down at 1000 rpm for 4 minutes. For sorting, cells were stained with 1 µg/ml Dapi (D1306, Invitrogen) in HF (2% FBS in HBSS (14175053, Gibco)), washed with HF and resuspended in HF. To determine the haploid population, a separate sample of cells was stained with 5 µM Vybrant® DyeCycleTM Violet Stain (V35003, Invitrogen) in full medium for 30 minutes and kept in vibrant containing medium. Cells were filtered with a 70 µm filter and then used for FACS sorting and analysis on a FACSARIA3 (BD, Franklin Lanes, NJ). Vybrant-stained cells were analyzed at 37° and used to set a size gate only containing haploid cells. Dapi-stained cells were single cell sorted at 4°C into 96-well plates, that were previously coated overnight with 0.1% gelatin (G9391, Sigma-Aldrich) in MQ and contained full medium supplemented with 1% penicillin/streptomycin (15140122, Gibco) and 0.025 M HEPES (H3375 Sigma-Aldrich, 1 M stock solution, pH 7.4, filter sterilized). The 3 independent clones used in this study were obtained from separate transfections of the same parental cell line. Resorting of the cell lines was performed with the same FACS procedure, with collection of cells in 15 mL tubes containing full medium with 1% penicillin and 0.025 M HEPES.
FACS data were analyzed and visualized with FlowJo™.

For qPCR analysis, 100,000 HAP1 cells per well were seeded on a 24-well plate. After 48 hours, cells were treated with indicated concentrations of CHIR99021. Cells were harvested 24 hours after treatment. RNA was isolated with Trizol (15596018, Invitrogen) according to the manufacturer’s instructions. cDNA was synthesized using SuperScriptIV (18090010, Invitrogen) according to the manufacturer’s instructions. qPCR was performed with SyberGreen (10710004, Invitrogen). The endogenous WNT target gene AXIN2 was amplified using primers RVA124 and RVA125, and HPRT housekeeping control was amplified using primers RVA24 and RVA25. Relative expression levels of AXIN2 were calculated using the comparative Delta-Ct method (Livak & Schmittgen, 2001; Schmittgen & Livak, 2008). Briefly, AXIN2 expression was normalized for HPRT expression and then the relative fold-change to a WT DMSO sample was calculated for all clones and conditions.

Luciferase Assay

For luciferase assays, 100,000 cells per well were seeded on a 24-well plate. Cells were transfected with 1µl X-tremeGene HP and 400 ng MegaTopflash reporter and 100 ng CMV-Renilla or 500 ng SGFP2-C1 as a negative control 24 hours later. Cells were treated with the indicated concentration of CHIR99021 24 hours after transfection and after another 24 hours medium was removed and the cells were harvested with 50 µl Passive Lysis Buffer (E1941, Promega). Luciferase activity was measured on a GloMax Navigator (Promega) using 10µl lysate in a black OptiPlate 96-well plate (6005279, Perkin Elmer, Waltham, MA) and 50 µL homemade firefly and luciferase reagents (according to (Fuerer, Nostro, & Constam, 2014; Hampf & Gossen, 2006)). Renilla and Luciferase luminescence values were corrected by
subtracting the background measured in the SGFP2-transfected control. MegaTopflash
activity was calculated as the ratio of corrected Firefly and Renilla luminescence and
normalized to the relative DMSO control.

**Western Blot**

The remaining lysates of the luciferase assay were cleared by centrifugation for 10 minutes at
12,000 g at 4°C. Western blot analysis was performed and quantified as previously described
(Jacobsen et al., 2016). Antibodies were used with the following dilutions, 1:1000 Non-
phosphorylated (Active) β-catenin clone D13A1 (8814S, Cell Signaling, Danvers, MA), 1:2000
total β-catenin clone 14 (610153, BD), 1:1000 α-Tubulin clone DM1A (T9026, Sigma-Aldrich),
1:1000 GFP polyclonal (A-6455, Invitrogen), 1:20,000 IRDye 680LT Goat anti-Rabbit IgG (926-68021, LI-COR, Lincoln, NE), 1:20,000 IRDye 800CW Donkey anti-Mouse IgG (926–32212, LI-
COR).

**Time-lapse imaging**

The day before imaging, 88,000 cells/well were seeded on an 8 well chamber slide with glass
bottom (80827-90, Ibidi, Gräfelfing, Germany). HAP1SGFP2-CTNNB1 clone 2 was used for the main
Figure 4, all 3 clones were used for Figure 4 supplement 1. Approximately 6 hours before
imaging, medium was replaced with full medium supplemented with 1%
penicillin/streptomycin, 0.025M HEPES and 500nM SiR-DNA (SC007, Spirochrome, Stein am
Rhein, Switzerland). Time lapse experiments were performed on an SP8 confocal microscope
(Leica Microsystems, Wetzlar, Germany) at 37°C with a HC PL APO CS2 63x/1.40 oil objective
(15506350, Leica), 488 and 633 lasers, appropriate AOBS settings, using HyD detectors for
fluorescent signal with a 496-555 for SGFP2-CTNNB1 and 643-764 bandpass for SiR-DNA, and
a transmission PMT. Using multi-position acquisition, up to 24 images were captured every 5
minutes. Focus was maintained using AFC autofocus control on demand for every time point and position. Automated cell segmentation and intensity quantification was performed using a custom CellProfiler™ pipeline (made available at https://osf.io/6pmwf/). Intensities were normalized per position to the average intensity in the cellular compartment (nucleus or cytoplasm) for that position before the addition of the compounds. The nuclear cytoplasmic ratio was calculated by dividing the raw nuclear intensity by the raw cytoplasmic intensity.

Movies and still images were extracted with FIJI/ImageJ.

**FCS and N&B cell preparation and general settings**

Two days before FCS and N&B experiments, 44,000 cells/well (HAP1\(^{WT}\) or HAP1\(^{SGFP2-CTNNB1}\) clone 2) were seeded on an 8-well chamber slide with a glass bottom (80827-90, Ibidi). The day before measuring, HAP1\(^{WT}\) cells were transfected with ~5 ng pSGFP2-C1, EGFP or EGFP dimer and ~200 ng pBlueScript KS(+) per well with X-tremeGene HP or Lipofectamine 3000. Lipofectamine 3000 yielded better transfection efficiency.

FCS and N&B measurements were performed on an Olympus FV-1000 equipped with SepiaII and PicoHarp 300 modules (Picquant, Berlin, Germany) at room temperature. An Olympus 60x water immersed UPLS Apochromat (N.A. 1.2) objective was used for FCS acquisition and Figure 3 supplement 1E, and an Olympus 60x silicon immersed UPLS Apochromat (N.A. 1.4) objective was used for N&B measurements. Green fluorophores were excited with a 488 nm diode laser (Picquant) pulsing at 20 MHz and detected through a 405/480-488/560/635 nm dichroic mirror (Chroma, Bellows Falls, VT) and 525df45 nm bandpass filter (Semrock, Rochester, NY) with an Avalanche Photodiode (APD) (MPD, Bolzano, Italy). For, figure 3 supplement 1E and for FCS and N&B reference images the same laser and dichroic were used,
but the signal was detected through a 505-540 bandpass filter with an internal PMT of the FV-1000 Olympus microscope.

**FCS data acquisition and analysis**

For FCS measurements, a confocal image of HAP1\(^{5GFP2-CTNNB1}\) was recorded. In this reference image, a single pixel was set as region of interest (ROI), based on its localization in the cytoplasm or nucleus as judged by the transmission image. In this ROI, the fluorescence intensity was measured over time using an APD, for typically 120 seconds.

FCS measurements were analyzed in FFS Dataprocessor version 2.3 (SSTC, Minsk, Belarus).

The autocorrelation curve \(G(\tau)\) was calculated from the measured intensity \(I\) according to equation 1. Intensity traces with significant photobleaching, cell movement or focal drift were excluded from further analysis. From other traces a portion of the trace with minimal (less than 10%) intensity drift or bleaching was selected to generate autocorrelation curve (AC).

\[
G(\tau) = 1 + \frac{\langle \delta I(t) * \delta I(t + \tau) \rangle}{\langle I \rangle^2} \tag{Eq. 1}
\]

The resulting AC was fitted with a Triplet-state-diffusion model, described in equation 2. \(G_{\infty}\) accounts for offset in the AC caused by intensity drift. \(N\) is the average of the number of particles that reside in the confocal volume. \(F_{\text{trip}}\) and \(\tau_{\text{trip}}\) describe the fraction of molecules in the dark state and the relaxation of this dark state respectively. Of note, in this case, \(F_{\text{trip}}\) and \(\tau_{\text{trip}}\) account both for blinking of the fluorescent molecules and for the afterpulsing artefact of the APD. \(\tau_{\text{diff,i}}\) describes the diffusion rate of the fluorescent molecules with the corresponding fraction, \(F_i\). This diffusion time depends on the structural parameter \(sp\), which is defined as the ratio of the axial \((\omega_z)\) over the radial axis \((\omega_{xy})\) of the observation volume.
\[ G(\tau) = G_\infty + \frac{1}{<N>} \left[ \frac{F_{\text{trip}}}{1 - F_{\text{trip}}} \right] e^{-\tau} \sum_i \left( 1 + \frac{\tau}{\tau_{\text{diff},i}} \right) \sqrt{1 + \frac{\tau}{\tau_{\text{diff},i} \cdot \text{sp}^2}} \]

Eq. 2

The apparent particle numbers \((N_{\text{apa}})\) for SGFP2-CTNNB1 were corrected for autofluorescence and bleaching (equation 3). The autofluorescence \((I_{\text{autofluorescence}})\) of HAP1 cells in the nucleus and cytoplasm was measured in untransfected HAP1 cells using the same settings as for FCS measurements. The correction for moderate bleaching is based on the intensity of the selected portion of the intensity trace for AC calculation \((I_{\text{ana}})\) and the intensity at the start of the measurement \((I_{\text{start}})\).

The size and shape of the observation volume was calibrated daily by measuring Alexa Fluor™ 488 NHS Ester (A20000, Molecular probes, Thermo Scientific, stock dilution in MQ) in PBS in a black glass-bottom cell imaging plate with 96 wells (0030741030, Eppendorf, Hamburg, Germany). From the FCS measurements of Alexa488, the \(\tau_{\text{diff}}\) and \(\text{sp}\) were determined by fitting with a single diffusion and blinking component. The diffusion coefficient \((D)\) of Alexa488 in aqueous solutions at 22.5 °C is 435 μm²s⁻¹ (Petrášek & Schwille, 2008). From these parameters, the axial diameter can be determined with equation 4 and the volume can be approximated by a cylinder (equation 5). This allows for transformation of particle numbers to concentrations (equation 5) and diffusion times to diffusion coefficients (equation 4) that are independent of measurement settings and small daily changes in alignment of the microscope.

\[ N_{\text{corr}} = N_{\text{apa}} \left[ 1 - \frac{I_{\text{autofluorescence}}}{I_{\text{total}}} \right]^2 \left[ \frac{I_{\text{start}}}{I_{\text{ana}}} \right] \]

Eq. 3
\[
\tau_{\text{diff}} = \frac{\omega_{xy}^2}{4D} \quad \text{Eq. 4}
\]

\[
V = 2\pi \omega_{xy}^3 \ast sp \quad \text{Eq. 5}
\]

\[
C = \frac{N_{\text{corr}}}{V * N_A} \quad \text{Eq. 6}
\]

The model to fit SGFP2-CTNNB1 measurements contained 2 diffusion components. The first diffusion component was fixed to the speed of monomeric SGFP2-CTNNB1. To estimate the speed of monomeric SGFP2-CTNNB1, the speed of free floating SGFP2, transfected in HAP1 cells, was measured to be 24.1 µm\(^2\)s\(^{-1}\) using FCS. Subsequently, this speed was used to calculate the speed of monomeric SGFP2-CTNNB1 with Einstein-Stokes formula (Equation 7).

As the temperature (T), dynamic viscosity (\(\eta\)) and Boltzmann’s constant (\(k_B\)) are equal between SGFP2 and SGFP2-CTNNB1 measurements, the expected difference in diffusion speed is only caused by the radius (\(r\)) of the diffusing molecule. The difference in radius was approximated by the cubic root of the ratio of the molecular weight of the SGFP2-CTNNB1 fusion protein (88 + 27=115 kDa) and the size of the SGFP2 protein (27 kDa), thus expecting a 1.62 times lower diffusion coefficient (compared to free floating SGFP2) of 14.9 µm\(^2\)s\(^{-1}\) for SGFP2-CTNNB1.

\[
D = \frac{k_B T}{6\pi \eta r} \quad \text{Eq. 7}
\]

In the fitting model, the structural parameter was fixed to the one determined by the Alexa488 measurements of that day. To ensure good fitting, limits were set for other parameters; \(G_{\text{m}}\) [0.5-1.5], \(N\) [0.001, 500], \(\tau_{\text{trip}}\) [1*10\(^{-6}\)-0.05 ms], \(\tau_{\text{diff2}}\) [10-200 ms].
N&B data acquisition and analysis

For N&B analysis, 50 images were acquired per measurement with a pixel time of 100 µs/pixel and a pixel size of 0.138-0.207 µm. The fluorescent signal was acquired with the APD described above for the FCS measurements. As a control and to optimize acquisition settings, HAP1 cells transfected with SGFP2, EGFP monomer or dimer were measured alongside HAP1SGFP2-CTNNB1 cells treated with BSA or WNT3A. APD readout was converted to a TIF stack using a custom build .ptu converter (Crosby et al., 2013). This TIF stack was further analyzed using an ImageJ macro script (modified from (Crosby et al., 2013), made available at https://osf.io/ys5qw/) based on Digman, Dalal, Horwitz, & Gratton, 2008.

Within the script, average brightness and particle numbers were calculated for nuclear or cytoplasmic ROIs, which were set based on transmission image. Static or slow-moving particles, including membrane regions, were excluded by thresholding and/or ROI selection, since they can severely impact the brightness measured.

Data representation and statistical analysis

Data processing and representation were performed in RStudio (version 1.1.456 running R 3.5.1 or 3.6.1). 95% confidence intervals of the median mentioned in the text and shown in Table 1-2 were calculated using PlotsOfDifferences (Goedhart, 2019). Representation of the imaging data in Figure 4 supplement 2 and in supplementary movies 3-4 were generated in RStudio using a script based on PlotsOfDifferences (made available at https://osf.io/sxakf/).

Model description

We developed a minimal model for WNT signaling based on a previous model from the Kirschner group (Lee et al., 2003). The R source code of the model is available at https://osf.io/2e8by/.
Our minimal model comprises the following reactions:

\[ CB + DC \overset{k_2}{\underset{k_1}{\rightleftharpoons}} CB^* - DC \]  
\[ CB^* - DC \overset{k_3}{\rightarrow} DC + CB^* \]  
\[ DVL + DC \overset{k_5}{\underset{k_4}{\rightleftharpoons}} DC^* \]  
\[ CB \overset{k_7}{\underset{k_6}{\rightarrow}} NB \]  
\[ NB + TCF \overset{k_9}{\underset{k_8}{\rightleftharpoons}} NB - TCF \]

1. Binding of cytoplasmic CTNNB1 (CB) to destruction complex
2. Release of phosphorylated CB (CB*) and recycling of the destruction complex
3. Inactivation of the destruction complex by DVL
4. Nucleocytoplasmic shuttling of CB to and from the nucleus
5. Binding of NB to TCF

Below, we show the differential equations that govern the concentrations of the different compounds over time for the reactions described above. Table 3 in the main text gives the correspondence between the variables (i.e. \(x_i\)) in the differential equations and the model name (i.e. CB) in the reactions. The parameter \(w\) in equations (7) and (8) is \(w = 0\) in the absence of WNT and \(w = 1\) if WNT is present, i.e. in our minimal model the inactive form of the destruction complex (DC*) is only present if WNT is present. The parameter \(b\) in equation (6) represents the constant production of CTNNB1, corresponding to \(v_{12}\) in Lee et al., 2003.

\[
\frac{dx_1}{dt} = -k_1x_1x_2 + k_2x_3 - k_6x_1 + k_7x_6 + b
\]

\[
\frac{dx_2}{dt} = -k_1x_1x_2 + (k_2 + k_3)x_3 - w(k_4x_2 - k_5x_4)
\]

\[
\frac{dx_3}{dt} = k_1x_1x_2 - (k_2 + k_3)x_3
\]

\[
\frac{dx_4}{dt} = w(k_4x_2 - k_5x_4)
\]

\[
\frac{dx_5}{dt} = k_6x_1 - k_7x_5 - k_8x_5x_6 + k_9x_7
\]

\[
\frac{dx_6}{dt} = -k_8x_5x_6 + k_9x_7
\]

\[
\frac{dx_7}{dt} = k_8x_5x_6 - k_9x_7
\]
Equilibrium conditions without WNT

The parameters in our model can in part be determined from our measurements of the equilibrium concentrations of CB, NB and their complexes, see Table 3-4 in the main text.

Where we could not determine the parameters from our measurements, we used published values as indicated.

Under equilibrium conditions, the concentrations of the compounds do not change with time and the left-hand side of equations (6) - (12) is zero. From equations (10) and (11) we can determine the ratio of the rate constants $k_6$ and $k_7$ from the measured values of $x_1$ and $x_5$:

$$k_6 x_1 = k_7 x_5 \iff \frac{k_6}{k_7} = \frac{x_5}{x_1} = \frac{95}{104} = 0.913$$  \hspace{1cm} (13)

From equations (6), (8), (10) and (11) we have:

$$-k_3 x_3 + b = 0 \iff k_3 = \frac{b}{x_3} = \frac{0.423}{67} = 0.0063$$  \hspace{1cm} (14)

Our reaction (1) corresponds closely to step 8 in Lee et al. therefore, we use the value of the dissociation constant $K_8 = 120$ nM from Lee et al. for our dissociation constant $K_1 = \frac{k_2}{k_1}$.

To calculate the dissociation constant for the NB-TCF complex, we estimate an equilibrium concentration for free TCF ($x_6$) from Tan et al. (2012). From their Figure 11 it is seen that the bound TCF concentration in equilibrium in the presence of WNT has about the same value as the initial free TCF concentration and that no initial bound TCF is present. However, we measured NB-TCF also in the initial state. Therefore, we consider the free TCF concentration value from Tan et al. as a lower bound for the estimate of total TCF. Also, from Figure 11 of Tan et al. (2012) we estimate that of the initial free TCF, a fifth remains in the nucleus as free TCF after WNT is turned on. We measured 102 nM NB-TCF in the nucleus after the application of WNT. This leads to an estimate of the total concentration of TCF, $TCF^0$, in the nucleus of:
\[ [TCF^0] = 102 + 0.2 \times 102 = 122 \text{ nM}. \] If we assume that the total TCF concentration does not change by the application of WNT, we calculate the dissociation constant of the NB-TCF complex from equation (12):

\[
k_8x_5(TCF^0 - x_7) = k_9x_7 \Rightarrow \frac{k_9}{k_8} = K_2 = \frac{x_5(TCF^0 - x_7)}{x_7} = \frac{95 \times 99}{23} = 409
\] (15)

**Equilibrium conditions with WNT**

We model the action of WNT by deactivation of the destruction complex by DVL through reaction 3 by setting \( w = 1 \) in equations (7) and (9). The dissociation constant of CB*-DC, \( K_1 \), is assumed not to change in the presence of WNT. The measurements of free CB and NB in equilibrium (see Table 2) give for the ratio of \( k_6 \) and \( k_7 \):

\[
k_6x_1 = k_7x_5 \leftrightarrow \frac{k_6}{k_7} = \frac{x_5}{x_1} = \frac{186}{145} = 1.28
\] (16)

The value of the rate of decay of the phosphorylated complex CB*-DC, \( k_3 \), is found to be the same for the "without WNT" situation:

\[-k_3x_3 + b = 0 \Leftrightarrow k_3 = \frac{b}{x_3} = \frac{0.423}{67} = 0.0063
\] (17)

To uniquely determine the ratio of \( k_4 \) and \( k_5 \), we need the concentrations of the destruction complex DC and DC+ neither of which we have access to. We can, however, fit this ratio with our model to the measured values of \( x_1 \) and \( x_7 \) and find \( k_4/k_5 = 2.56 \).

We again calculate the dissociation constant of the NB-TCF complex from equation (12), using the concentrations for NB and NB-TCF obtained with FCS.

\[
k_8x_5(TCF^0 - x_7) = k_9x_7 \Rightarrow \frac{k_9}{k_8} = K_2 = \frac{x_5(TCF^0 - x_7)}{x_7} = \frac{186 \times 20}{102} = 36.5
\] (18)
Notice that we determined the ratios of the rate constants from the measured equilibrium values of free and bound CTNNB1 in the cytoplasm and the nucleus. This means that our rate constants are determined up to a multiplicative factor: the equilibrium equations do not change if all rate constants $k_i$ and the parameter $b$ are multiplied by the same factor, $R$. The factor $R$ determines the rate at which our model system reaches equilibrium. By comparing the times equilibrium was reached by the cytoplasmic and nuclear SGFP2-CTNNB1 signals (Figure 4 C, D) of about 4.5 hours, we fitted a factor $R = 20$ for our model.
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Competing interest statement

The authors have no competing interests to declare.
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Supplementary Movie legends

Supplementary Movie 1-3: Representative movies of confocal time-lapse experiments, showing SGFP2-CTNNB1 (left, green), SiR-DNA staining (middle, magenta) and transmission image (right, grey) after treatment with vehicle control (BSA) (Supplementary Movie 1), 100 ng/ml WNT3A (Supplementary Movie 2) or 8 µM CHIR99021 (Supplementary Movie 3). Time of addition is at 00:00:00 (indicated at the top left). Scale bar in the lower right represents 20μm.

Supplementary Movie 4-6: Movies showing the quantification of time-lapse microscopy series (from Figure 4 and Supplementary Movie 1-3) at each time point showing all individual cells from 3 biological experiments. Time of addition of the indicated substances is at 00:00:00 (indicated at the top left). The left graph represents the raw data (colored dots, each dot is one cell, n=155-400 cells for each condition and time point), the median (black circle) and the 95% CI of the median (black bar). The right graph represents the median difference (black circle) from the treatments to the control (BSA). When the 95% CI (black bar) does not overlap 0, the difference between the two conditions is significant.

Supplementary Movie 4: Quantification of the normalized intensity of SGFP2-CTNNB1 in the cytoplasm. Significant changes in intensity can first be observed after 40 minutes of 8 µM CHIR99021, and after 70-80 minutes of 4 µM CHIR99021 or 25-100ng/ml WNT3A treatment.

Supplementary Movie 5: Quantification of the normalized intensity of SGFP2-CTNNB1 in the nucleus. Significant changes in intensity can be observed for all treatments (but not controls) after 20-50 minutes.

Supplementary Movie 6: Quantification of the nuclear-cytoplasmic ratio of SGFP2-CTNNB1, calculated from raw intensity values underlying Supplementary Movies 4 and 5. Significant changes in the nuclear-cytoplasmic ratio can be observed for all treatments (but not controls) after 20-50 minutes.