ROS-induced near-homozygous genomes in thyroid cancer

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Abstract

A near-homozygous genome (NHG) is especially seen in a subset of follicular thyroid cancer of the oncocytic type (FTC-OV). An NHG was also observed in the metabolically relatively quiescent cell lines XTC.UC1, a model for FTC-OV, and in FTC-133, -236 and -238, the latter three derived from one single patient with follicular thyroid cancer. FTC-236 subclones showed subtle whole-chromosome differences indicative of sustained reciprocal mitotic missegregations. Reactive oxygen species (ROS) scavenger experiments reduced the number of chromosomal missegregations in XTC.UC1 and FTC-236, while pCHK2 was downregulated in these cells. Treatment with antimycin A increased ROS indicated by enhanced MitoSOX Red and pCHK2 fluorescence in metaphase cells. In a selected set of oncocytic follicular thyroid tumors, increasing numbers of whole-chromosome losses were observed toward an aggressive phenotype, but with retention of chromosome 7. Together, ROS activates CHK2 and links to the stepwise loss of whole chromosomes during tumor progression in these lesions. We postulate that sequential loss of whole chromosomes is a dominant driver of the oncogenesis of a subset of follicular thyroid tumors.

Key Words
- thyroid neoplasm
- oncocytic
- reactive oxygen species
- chromosome instability
- checkpoint kinase 2

Introduction

Follicular thyroid cancer of the oncocytic variant (FTC-OV) is overrepresented in recurrent non-medullary thyroid cancer (NMTC) (Wada et al. 2002). FTC-OV is characterized by a strong eosinophilic cytoplasm due to abundant mitochondria in conjunction with a reprogrammed metabolism probably driven by estrogen-related receptor α (Mirebeau-Prunier et al. 2013). In FTC-OV mitochondrial DNA (mtDNA), variants in complexes of the respiratory chain are also found, which might cause a restricted ATP production (Bonora et al. 2006). Nonetheless, disruptive and/or damaging mtDNA variants are not exclusive to oncocytic tumors, since these mutations were as frequently found in non-oncocytic papillary thyroid and other cancers (Chatterjee et al. 2006, Zhou et al. 2007, Larman et al. 2012, Ju et al. 2014).

FTC-OV has been extensively studied at the genomic level (Maximo et al. 2014). Based upon flow or image DNA cytometry, comparative genomic hybridization (CGH) experiments or interphase fluorescence in situ hybridization, these tumors were considered diploid or aneuploid with whole-chromosome gains (McLeod et al. 1988, Flint & Lloyd 1990, Salmon et al. 1993, Roque et al. 1999, Tallini et al. 1999, Erickson et al. 2001, Wada et al. 2002, Dettori et al. 2003). Using a new method for DNA content analysis (Corver et al. 2005) in combination with single nucleotide polymorphism (SNP) technology

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(Corver et al. 2008), we demonstrated that many FTC-OV actually show a near-homozygous genome (NHG) in which a phase of near-haploidization is followed by endoreduplication or genome doubling of the entire NHG (Corver et al. 2012, 2014). Our observations regarding an NHG have been confirmed by others (Wagle et al. 2014, Kasaian et al. 2015). Furthermore, retention of chromosome 7, a hallmark for FTC-OV, was recently attributed to genomic imprinting of genes on both paternal and maternal alleles that are important for tumor cell survival (Boot et al. 2016).

The loss of many entire chromosomes (whole-chromosome instability, w-CIN) as found in tumors with an NHG suggests chromosome missegregations or non-disjunctions during mitosis. Despite the fact that molecular mechanisms implicated in w-CIN are studied extensively, a common cause in cancer has so far not been found.

Recently, Bakhoum and coworkers demonstrated that doxorubicin or ionizing radiation (IR) during mitosis induced a DNA damage response with lagging chromosomes (Bakhoum et al. 2014). Exposing cells to chloroquine, an activator of ATM-kinase, induced lagging chromosomes to the same levels but without significant DNA damage. ATM can also be activated by oxidative stress in the absence of DNA double-strand breaks (Alexander et al. 2010, Guo et al. 2010). These observations point to a concomitance between an altered metabolism and w-CIN in a subset of thyroid cancer cells.

Here, we hypothesized that increased levels of reactive oxygen species (ROS) underlie the process of w-CIN in thyroid tumors with NHG. We used DNA content analysis and SNP array technology to search for an NHG in thyroid cancer cell lines in order to investigate the underlying processes. We identified XTC. UC1 cells, the only known cell line model for FTC-OV, and the follicular thyroid cancer cell lines FTC-133, FTC-236 and FTC-238 as models for NHG. Next, we studied the response to metabolic stress and determined the role of ROS and CHK2 activation in whole-chromosome segregation errors by pharmacologic intervention using high-resolution immunofluorescence microscopy as a read-out system in NHG (XT. UC1, FTC-236) and non-NHG (BHP 2–7, SW579) thyroid cancer cell lines. Finally, we verified our in vitro findings in clinical samples and searched for the sequence of whole-chromosome gains or losses in the progression of oncocytic thyroid follicular adenoma to carcinoma.

### Methods

#### Patient material

Selection of cases: SNP array and flow cytometric DNA content data were collected from a cohort of 88 recurrent NMTC cases. Of these, 33 cases showed oncocytic histology by routine H&E histology, determined by an experienced pathologist (HM). Of these, 25 cases showed clear marks of w-CIN (range 1–21 whole-autosomes). One sample was a FTC-OV with the loss of entire chromosome 22. The remaining 24 cases all showed a follicular phenotype and were used to construct a model for the sequential loss of whole chromosomes: 7 oncocytic follicular thyroid adenomas (FA-OV), 1 minimal-invasive oncocytic thyroid follicular carcinoma (FTC-OV-MI), 14 more advanced FTC-OV and 2 anaplastic thyroid carcinomas (ATC) that derived from FTC-OV were included in this study (data submitted to Gene Expression Omnibus, [https://www.ncbi.nlm.nih.gov/geo](https://www.ncbi.nlm.nih.gov/geo)). Morphological evaluation (H&E sections) was performed on the primary tumors without exception. From three patients (Study Nos. 8, 17 and 21), only a metastasis was available for further analysis. From all other patients, primary tumor material was used. Additional patient information on gender, age of diagnosis, invasiveness, recurrence and death of disease can be found in Supplementary Table 1, see section on supplementary data given at the end of this article.

From one patient (No. 15), both formalin-fixed, paraffin-embedded (FFPE) tissue and frozen tissue were available. The oncocytic phenotype showed to be highly associated with whole-chromosomal aberrations ($P<0.0001$, Fisher’s exact, two-sided). H&E sections from representative adenoma and cancer cases (Nos. 1, 3, 5, 7, 8, 18 and 23) were scanned with a high-resolution imager (40× magnification, 0.25 µm/pixel, Philips Ultra Fast Scanner 1.6 RA, Philips). Sections (2× and 40× digital zoom) are shown in Supplementary Fig. 1A (Nos. 1, 3, 5 and 7) and B (Nos. 8, 18 and 23).

From the 55 non-oncocytic cases, 6 showed w-CIN (range 1–12 whole-autosomes) and 49 cases showed structural/segmental chromosomal instability (s-CIN, samples show segmental chromosomal defects: chromosomal arms or segments are lost or amplified), a combination of s-CIN and w-CIN or no chromosomal aberrations. One FTC-OV showed s-CIN only. All samples are summarized in Table 1. Samples were handled according to the medical ethical guidelines as described in the Code Proper Secondary Use of Human Tissue (Dutch Federation of Medical Sciences, [www.federa.org](http://www.federa.org)).
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Cell lines
The following cell lines were used: BHP 2–7 (derived from a primary papillary thyroid cancer) (Ohta et al. 1997), FTC-133, FTC-236, FTC-238 (derived from three different metastases of an FTC) (Demeure et al. 1992, Goretzki et al. 1990), SW579 (deposited by A Leibovitz) and TT2609-C02 (Geldof et al. 2001), all derived from an FTC and XTC.UC1, the only official model for FTC-OV (Zielke et al. 1998). All cell lines were cultured under standard conditions (37°C, 5% CO₂ and 95% humidified air). Media were supplemented with glutamine, 10% heat-inactivated FCS and 50 U/mL penicillin and 50 µg/mL streptomycin. FTC-133, FTC-236, FTC-238 and XTC.UC1 were cultured in DMEM-F12. SW579 and TT2609-C02 (Geldof et al. 2001), all derived from an FTC and XTC.UC1, the only official model for FTC-OV (Zielke et al. 1998). All cell lines were cultured under standard conditions (37°C, 5% CO₂ and 95% humidified air). Media were supplemented with glutamine, 10% heat-inactivated FSC and 50 µg/mL streptomycin. FTC-133, FTC-236, FTC-238 and XTC.UC1 were cultured in DMEM-F12. SW579 and TT2609-C02 were maintained in RPMI medium. HeLa cells were taken along as control in metabolic testing. A short tandem-repeat multiplex PCR (Cell ID GenePrint 10 system, Promega) authenticated the cell lines which were no longer cultured than 20 passages and were mycoplasma tested biweekly (in-house PCR method). From FTC-236, seven subclones were generated by limiting dilution: clones A–C, E and G–I. These subclones were maintained in complete DMEM-F12 medium and are available in our laboratory.

DNA content analysis by flow cytometry
For DNA content analysis on cell lines, the Vindeløv method was used (Vindelov et al. 1983). In short, cells were harvested by trypsin/EDTA (Corver et al. 1995), counted and 2 × 10⁶ cells were divided over two polystyrene tubes. Cells were pelleted and to one tube, trout red blood cells (TRBC) were added as internal reference for DNA content. A short tandem-repeat multiplex PCR (Cell ID GenePrint 10 system, Promega) authenticated the cell lines which were no longer cultured than 20 passages and were mycoplasma tested biweekly (in-house PCR method). From FTC-236, seven subclones were generated by limiting dilution: clones A–C, E and G–I. These subclones were maintained in complete DMEM-F12 medium and are available in our laboratory.

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DNA isolation
For SNP array analysis (iCOG and HumanCytoSNP-12, Illumina, Inc., San Diego, CA, USA) and mtDNA mutation analysis, high-molecular weight DNA was extracted from frozen samples or frozen cell lines pellets, respectively.

For SNP array analysis of FFPE tissues (GoldenGate assay, Illumina), three 0.6-mm diameter tissue punches (Beecher Instruments, USA) were taken from selected tumor areas (HM) in order to enrich for tumor DNA. Next, tissue punches were dewaxed prior to DNA isolation, and overnight digested with proteinase K at 56°C.

DNA from frozen and FFPE samples was extracted using the NucleoSpin purification kit (Macherey-Nagel GmbH & Co. KG, Düren, BRD, Germany) according to the manufacturer’s instructions. DNA concentrations were determined using the Picogreen method (Thermo Fisher Scientific).

Genome-wide SNP array analysis
HumanCytoSNP-12 BeadChip incorporates over 220,000 SNPs across the genome. The iCOG arrays are custom Illumina iSelect genotyping arrays that comprise over 200,000 SNPs. The two high-density platforms were evaluated in-house and showed similar results.

Table 1  Overview of structural and/or whole-chromosome aberrations found in 88 NMTC cases.

<table>
<thead>
<tr>
<th>Histology</th>
<th>Chromosomal aberration non-oncocytic</th>
<th>Chromosomal aberration oncotypic variant (OV)</th>
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<tr>
<td></td>
<td>s-CIN</td>
<td>w-CIN</td>
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<tr>
<td>FA</td>
<td>2</td>
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<td>FTC-MI</td>
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<tr>
<td>FTC</td>
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<td>FTC</td>
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<tr>
<td>ATC</td>
<td>1</td>
<td>1</td>
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<tr>
<td>Total</td>
<td>16</td>
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</table>

*Samples with an NHG with loss of chromosomes 1–4, 6 and 11 as a minimum genomic signature of FTC-OV; **1/7 FA showed an NHG.

ATC, anaplastic thyroid carcinoma; FA, follicular adenoma; FTC, follicular thyroid carcinoma; FTC-MI, follicular thyroid carcinoma, minimal invasive; FTPC, follicular variant papillary thyroid carcinoma; NC, no changes observed after LAIR analysis; PTC, papillary thyroid carcinoma; s-CIN, structural chromosomal instability; s/w-CIN, combination of s-CIN and w-CIN; w-CIN, structural chromosomal instability.
Arrays were run by a genomics service provider (ServiceXS, Leiden, Netherlands) and hybridized according to the manufacturers’ recommendations. The arrays were analyzed as described previously (Corver et al. 2008), using the beadarraySNP package (Oosting et al. 2007). The GoldenGate is a low-density platform but suitable for fragmented DNA from FFPE tissues. During analysis, the DNA index was incorporated allowing reliable copy number estimates. The method is referred to as Lesser-allele intensity-ratio (LAIR) analysis (Corver et al. 2008).

**mtDNA sequencing of cell line DNA**

Sanger sequencing of mtDNA was performed as extensively described previously (Corver et al. 2014). In short, 24 primer pairs (Eurofins MWG Operon, Ebersberg, Germany) were used covering the entire mitochondria and blasted against the National Centre for Biotechnology Information mitochondrial reference genome NC_012920 giving a 100% match. M13 tails were added to the primers for universal sequencing. Purified PCR products were Sanger-sequenced in both directions at MacroGen (Amsterdam, Netherlands). Data were analyzed using Mutation Surveyor version 4.0.8 (Softgenetics, State College, PA, USA). Variants were called against a GenBank reference (NC_012920) and checked manually. Next, coding missense mutations were selected and analyzed in silico using the Polyphen V2 package (Adzhubei et al. 2013).

**Inhibitors and small molecules**

Chemicals and small molecules used: carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP) (Cayman Chemical Company), range 0.5–2.0µM (stock 30mM in 100% dimethylsulfoxide (DMSO)); Buthionine sulfoximine (BSO), range 0.01–10mM ((stock solution 300 in HBSS); M36008, Thermo Fisher Scientific). Next, cells were permeabilized with PBS/1%BSA/0.5% Tween20 after fixation (4% PFA), and indirectly stained for pCHK2 (Thr68) (rabbit IgG, Cat. No. 2197, Cell Signaling Technology).


**Immunofluorescence and imaging**

Cells were cultured on alcohol-sterilized APES-coated glass slides (Thermo Fisher Scientific) or in 96-well µClear plates (Cat. No. 655090, Greiner Bio-One International GmbH, Austria), fixed with 4% phosphate-buffered paraformaldehyde (PFA) at 37°C and permeabilized with 100% ice-cold methanol and/or PBS/1%BSA/0.5% Tween20, also used as a blocking step. Staining: α-tubulin and HEC1 followed by a mixture of goat anti-mouse IgG1-AF488 or IgG2a-AF594 supplemented with 0.5µM Hoechst33342 (Sigma).

Oxygen anion (O₂⁻) detection: cells were treated with 2µM antimycin A for 30 min or left untreated, washed and incubated for 15 min with MitoSox red (Cat. No. M36008, Thermo Fisher Scientific). Next, cells were permeabilized with PBS/1%BSA/0.5% Tween20 after fixation (4% PFA), and indirectly stained for pCHK2 (1:200) and DNA (Hoechst33342, 0.5µM). Cells were studied using a DM6B fluorescence microscope (Leica). Images were acquired using a CoolPix camera and a 40× immersion oil objective (glass slides) or 40× dry objective (µClear plates).

**Immunoblotting**

Odyssey nitrocellulose membranes (LI-COR Biosciences) were washed and blocked in TBS Odyssey Blocking Buffer (LI-COR Biosciences) and incubated with an anti-p-CHK2-Thr68/anti α-tubulin mixture. A secondary mixture (GaRlgG-800CW and GaMIgG-680RD) was used for 30 min.
at room temperature. Protein bands were visualized using the Odyssey Classic scanning system.

**Oxygen consumption and glycolysis**

Twenty thousand cells of each cell line were plated in triplicate in a Seahorse XF-96 microwell plate and cultured overnight. TT2609-C02 did not attach well to the polystyrene and was left out. The cervical carcinoma cell line HeLa was included as a reference (Lee et al. 2015). Cells were challenged with different concentrations (0.5–2.0 µM, with steps of 0.5 µM) of FCCP according to the manufacturer instructions or left untreated. Oxygen consumption rates (OCR) and extracellular acidification rates (ECAR) were determined using a Seahorse XF-96 extracellular flux analyzer (Agilent). The OCR and ECAR data were combined into an energy profile.

**Statistical analysis**

Statistics for lagging, bridged or acentric chromosomes were calculated with a two-tailed Fisher's exact test using Prism 6 software (GraphPad).

**Results**

**w-CIN in thyroid cancer cell lines leads to an NHG**

Recently, we proposed a progression model for FTC-OV involving the stepwise loss of entire chromosomes during cancer progression finalizing in an NHG (Corver et al. 2012). To find evidence for our hypothesis, we studied seven NMTC cell lines for whole-chromosome losses by the LAIR analysis (Corver et al. 2008). XTC.UC1 clearly showed an NHG with most chromosomes in a pure homozygous state ((AA) or (AAA)) (Fig. 1A). The NHG profile in XTC.UC1 was expected, since a CGH copy number profile showed near-identical NHGs but differed in morphology and chromosome copy numbers as observed by the DNA index (Supplementary Fig. 3). From these observations, we deduced a model of segregation errors of chromosome 18 in FTC-236 (Fig. 1D). Alternatively, the missegregated chromosome may lag behind and forms a micronucleus after mitosis. Indeed, FTC-236 showed approximately 10 times higher numbers of micronuclei than BHP 2–7 under standard culture conditions (Supplementary Fig. 4). The LAIR profiles of BHP 2–7 and SW579 were clearly different and showed s-CIN (BHP 2–7) or a mixture of w-CIN and s-CIN (SW579) (Supplementary Fig. 5). TT2609-C02 was published earlier and also showed a mixture of w-CIN and s-CIN (Boot et al. 2016).

Collectively, these in vitro findings further support our proposed hypothesis that a subset of follicular thyroid cancers, especially those with oncocyctic metaplasia, show sustained missegregations during mitosis with selection pressure on whole-chromosome losses leading to NHG.

**FTC cell lines harboring an NHG show a quiescent metabolic phenotype**

FTC-OV is tightly associated with an altered metabolism (Mirebeau-Prunier et al. 2013). XTC.UC1 is a model for clinical FTC-OV and shows a relative quiescent glycolysis and oxidative metabolism (Lee et al. 2015). Since FTC-133, -236 and -238 also harbor an NHG, we questioned whether these cells show a comparable metabolism. The energy profiles of FTC-133, FTC-236 and FTC-238 were highly comparable to the relative quiescent energy profile of XTC.UC1 (Fig. 2A). In contrast, BHP 2-7, SW579 and HeLa showed a higher basic ECAR and OCR after exposure to FCCP.

Owing the highly comparable energy profiles, we next questioned whether FTC-133, -236 and -238 also harbor explanatory disruptive mtDNA mutations like XTC.UC1. We sequenced the entire mitochondrial genomes of the seven NMTC cell lines used in this study using twenty-four overlapping fragments (Corver et al. 2014). We confirmed the known disruptive C insertion at bp3571, causing a stop
codon at amino acid 101 (\textit{MT-ND1}, subunit of complex I) and the damaging mutation (m.15557 G > A: p.271E > K) in \textit{MT-CYB} (complex III) (Bonora \textit{et al.} 2006) in XTC.UC1. No further disruptive mutations were found in the remaining cell lines. Only damaging mtDNA mutations and silent polymorphisms were observed (Fig. 2B and Table 2). This supports our earlier observation in clinical samples that mtDNA mutations neither associate with an NHG nor with a relative quiescent energy profile (Corver \textit{et al.} 2014).

In conclusion, a modified relatively quiescent metabolic phenotype might be linked to the establishment...
of an NHG in FTC-OV. In the tested thyroid cancer cell lines, a relation between the mtDNA status and a certain metabolic phenotype or an NHG is not evident.

**ROS is linked to chromosome missegregation errors in FTC-OV**

Next, we questioned how an altered metabolism is related to an NHG in the NMTC cell line models FTC-133, -236, - 238 and XTC.UC1. XTC.UC1 is known to generate high levels of ROS (Stankov et al. 2006) and complex III, which plays a significant role in regulating ROS (Chen et al. 2003), is overexpressed in FTC-OV (Baris et al. 2005). Since ROS can activate ATM-CHK2 in a DNA damage-independent manner (Alexander et al. 2010, Guo et al. 2010) and ATM-CHK2 activation can cause lagging chromosomes (Bakhoum et al. 2014), ROS might underlie the development of an NHG.

Under basal conditions, XTC.UC1 cells and FTC-236 cells showed significantly higher numbers of lagging chromosomes than SW579 and BHP 2-7 cells ($P = 0.006$), whereas XTC.UC1 and FTC-236 were highly comparable ($P > 0.5$). Lagging chromosomes but also chromatin bridges, acentric chromatin or combinations were primarily found in XTC.UC1 and FTC-236 (Fig. 3A). XTC.UC1 showed significantly more chromosome bridges ($P = 0.0002$) and acentric chromosomes than BHP 2–7 ($P = 0.027$).

We then asked whether we can reduce the number of segregations errors in the latter two cell lines using pharmacological intervention affecting ROS. BHP 2–7 was taken for comparison. NAC is a potent ROS scavenger and serves as a precursor of cysteine for glutathione (GSH) synthesis, an important cellular antioxidant (Samuni et al. 2013). Treating the cells with 5–15 mM NAC significantly decreased the number of anaphase missegregations ($P < 0.05$) in XTC.UC1 and FTC-236 compared to high basal levels (Fig. 3B).

Next, we questioned whether CHK2 might be involved. Indeed, XTC.UC1 cells, the oncocytic cell line with a high frequency of spontaneous chromosome missegregations, showed phosphorylation CHK2 under standard culture conditions. pCHK2 diminished after glutathione addition in a dose-dependent manner. Treating the cells with 5–15 mM NAC significantly decreased the number of anaphase missegregations ($P < 0.05$) in XTC.UC1 and FTC-236 compared to high basal levels (Fig. 3B).

Next, we questioned whether CHK2 might be involved. Indeed, XTC.UC1 cells, the oncocytic cell line with a high frequency of spontaneous chromosome missegregations, showed phosphorylation CHK2 under standard culture conditions. pCHK2 diminished after glutathione addition in a dose-dependent manner. Treating the cells with $H_2O_2$ significantly increased the signal amplitude of CHK2 phosphorylation (Fig. 3C).

We then induced superoxide anion ($O_2^-$), the main component of mitochondrial ROS, by inhibiting complex III using antimycin A. A moderate increase in CHK2 phosphorylation was observed (Fig. 3C) exclusively caused by further CHK2 activation in metaphase nuclei (Fig. 3D).
which simultaneously showed an increase in MitoSOX Red fluorescence, an $\text{O}_2^-$ indicator (Fig. 3D). Interphase nuclei did not show pCHK2 fluorescence. The same was observed in BHP 2–7 cells but with lower fluorescent signal amplitudes (Supplementary Fig. 6). By treating BHP 2–7 cells with BSO, an inhibitor of $\gamma$-glutamylcysteine synthetase, the rate-limiting enzyme for GSH synthesis, the number of missegregations increased significantly and a subtle increase in CHK2 phosphorylation were observed in mitotic cells (Supplementary Fig. 7). This demonstrated that the ATM-CHK2 pathway is active in these cell lines and that $\text{O}_2^-$ can regulate the levels of pCHK2. Thus, ROS is likely involved in chromosomal missegregations in XTC.UC1 and in FTC-236 thyroid cancer cells by activating CHK2 signaling.

**Whole-chromosome segregation errors underlie the progression toward NHG**

To also find evidence for a sustained process of whole-chromosome reciprocal missegregations in (oncocytic) follicular thyroid tumors cases, we compared 7 FA-OV, 15 FTC-OV and 2 ATC both derived from an FTC-OV.

All carcinoma cases and 3 FA-OV in this study showed whole-chromosome losses. Strikingly, two FA-OV and the FTC-OV-MI case presented imbalances of whole chromosomes which showed homozygous in 100% (chromosomes 1 and 2), 94% (chromosomes 8 and 22), 88% (chromosome 9) and 65% (chromosome 18) of the FTC-OV samples ($n=17$) (Fig. 4A), indicating that these losses might be early events in tumor development. Additional losses of chromosomes 2, 3, 6, 11, 14–16 and 21 seem to be associated with tumor progression with losses of chromosomes 1–4, 6 and 11 as minimum signature of FTC-OV. Four FA-OV showed copy number gains only.

From one FTC-OV, we observed differences between two samples (Nos. 15a and 15b). Sample 15a showed heterozygous (AABB) for chromosomes 12, 13 and 18, while sample 15b is homozygous for these chromosomes (AA). This clearly demonstrates intra-tumor heterogeneity which can be the result of a sustained process of w-CIN active in FTC-OV cases. Whole-genome DNA sequencing (Nos. 11, 15a and 23) further confirmed our SNP array findings (Supplementary Fig. 8).

Together, our observations point to an initiating asymmetric mitotic event in which the retention of chromosome 7 (status (AAB) or (AB)), and, to a lesser extent, chromosomes 5 and 12 is crucial and selected during further progression. Apparently, the reciprocal counterparts with chromosomes 5, 7 and/or 12 in a homozygous state (A) do not survive or favor a selective outgrowth, likely driven...
by imprinted genes on chromosome 7 (Boot et al. 2016).
Genomic ranking further supports our hypothesis of a sustained process of whole-chromosome segregation errors and selection for whole-chromosome losses (Fig. 4B) in the progression from FA-OV to FTC-OV and ATC-derived FTC-OV. Progression by additional chromosomal losses (Study Nos. 13–24, Fig. 4B) seems to be associated with death of disease (Supplementary Table 1).

**Figure 3**
ROS is linked to missegregations via CHK2 activation. (A) Examples of spontaneous missegregations in XTC.UC1 showing: a lagging chromosome (upper row), a chromatin bridge (middle row) and an acentric chromosome (lower row). From left to right: kinetochore (red, AF594 fluorescence), α-tubulin (green, AF488 fluorescence), DNA (cyan, Hoechst33342 fluorescence), merged image. All images were captured with a 40× dry objective. An identical digital zoom was applied to all images. (B) XTC.UC1 and FTC-236 show a significant higher number of lagging chromosomes than BHP 2–7 and SW579 (P=0.0004) (n=300). Strong reduction (P<0.0001) of the percentage of mainly lagging chromosomes in XTC.UC1, FTC-236 and BHP 2–7 after treatment the cells for 2h with the ROS scavenger NAC. Black: lagging chromosomes, gray: chromosome bridge, white: acentric chromosomes. (C) Western blot showing auto-phosphorylation of CHK2 in XTC.UC1 cells (in green), α-tubulin internal control (red). CHK2 can be further activated by H2O2, while glutathione treatment reduced the amplitude of phosphorylation. An increase in pCHK2 was shown after incubating the cells with antimycin A. (D) XTC.UC1 cells were treated with antimycin A (lower panels) or left untreated (upper panels) and stained for ROS, pCHK2 and DNA. Increased ROS activity (MitoSOX Red, in green) was observed in many metaphase cells (see red arrows), which coincided with an increased pCHK2 fluorescence (in yellow, AF647 fluorescence). DNA was stained with Hoechst33342 (cyan). All images were captured with a 20× dry objective. An identical digital zoom was applied to all images.

**Discussion**
We propose that ROS is linked to the massive loss of whole chromosomes during oncocytic follicular thyroid tumor progression resulting in an NHG. Despite the fact that molecular mechanisms implicated in w-CIN are studied extensively; a common cause leading to near-haploidy in cancer is yet to be identified. Even in the era of
**Figure 4**
Genomic ranking and proposed tumor progression model for oncocytic follicular thyroid cancer. (A) Ranking of oncocytic follicular thyroid tumors based on whole chromosome losses or gains. The DNA index was obtained by multiparameter flow cytometry of cell suspension derived from FFPE tissue punches. Seven samples were bimodal (B) and samples 3, 10, 15, 19, 20, 23 and 24 show endoreduplication (genome doubling). (B) Sum of the relative allelic score showing progressive whole-chromosome losses or gains. Note the relative sharp transition from FA-OV to FTC-OV. (C) Newly proposed model for FTC-OV tumor progression based on the activation of CHK2 via direct ATM oxidation by high ROS. This results in increased tension of the kinetochore/α-tubulin network leading to whole-chromosome segregation errors (Bakhoum et al. 2014). The progression from FA-OV seems to be driven by a stepwise loss of whole chromosomes, but with retention of chromosome 7 due to maternal- and paternal-imprinted genes important for survival (Boot et al. 2016). The loss of chromosome 22 is an early event. Blue, heterozygous (AB) or (AABB), score 2.0; light blue, imbalance (AAB), score 3.0; purple, imbalance composed of a mixture of status (AB) and (A), score 1.5; red, homozygous (A) or (AA), etc., score 1.0; B, bimodal DNA histogram (underlined: major population); U, unimodal DNA histogram.
next-generation sequencing (NGS), candidate gene variants directly related to whole-chromosome segregation errors are rarely found. For example, inactivation of the long non-coding RNA NORAD results in chromosome bridges and mitotic slippage, a common route to high aneuploidy (Lee et al. 2016). However, chromosome bridges do not explain the whole-chromosome losses found in FTC-OV with NHG. Relative low numbers of chromosomal bridges were found in XTC.UC1 and FTC-236. Moreover, in FTC-OV, gene variants or fusions in cancer driver genes are relatively rare, although those found (e.g. FLCN, MEN1, mTOR, PTEN, PIK3CA, TP53 and TSC2, Table 3) are frequently involved in metabolic switches. Possibly additional tumor drivers need to be identified. Indeed, we only found a PTEN- and a TP53-inactivating gene variant after whole-genome sequencing (WGS) of three FTC-OV with NHG (data not shown). Thus, FTC-OV does not seem to share the same mutation(s) or share the same mutated gene(s) which can be directly related to w-CIN.

Alternatively, we now suggest that the ubiquitous loss of whole chromosomes in FTC-OV with NHG might be a tumor driver induced by oxidative stress. ROS are the main mitochondrial side products. As mentioned earlier, ROS can activate CHK2 in a DNA damage-independent manner (Alexander et al. 2010, Guo et al. 2010), while CHK2 activation during mitosis induces chromosome segregation errors (Bakhoum et al. 2014).

By pharmacological intervention using the antioxidant NAC, we were able to significantly reduce the ongoing process of chromosome missegregations in XTC.UC1 and FTC-236 cells, models for NHG-associated oncocytic – and follicular thyroid cancer. The missegregations are probably driven by CHK2 since glutathione treatment diminished CHK2 autophosphorylation.

Our work also implies that the process of whole-chromosome segregation errors is on-going in clinical FTC-OV, demonstrated by intra-tumor heterogeneity (samples 15a and 15b) and the progressive loss of whole chromosomes toward subsets of ATC that is derived from FTC-OV (samples 23 and 24) with only chromosomes 7 and 12, and 7 in a heterozygous state, respectively. The latter observation was confirmed by others in a particular case of an ATC with responsiveness to everolimus (Wagle et al. 2014). Furthermore, progressive loss of whole chromosomes seems to be associated with a poor prognosis. Especially, the thyroid cancer patients with extensive whole-chromosome losses (twelve out of seventeen) died of disease (Supplementary Table 1), though this must be confirmed in a larger cohort. Using FTC-236 as an in vitro model, we showed that sister chromatids can be asymmetrically distributed over the two sister cells after a reciprocal mitotic event and that both sister cells are viable as demonstrated by the outgrowth of these clones. However, genomically reciprocal tumors were not observed in our cohort of clinical samples. Markedly, chromosomes 1–4, 6 and 11, which mainly comprises the larger chromosomes, were preferentially lost in FTC-OV, whereas overall in human carcinomas, the smaller chromosomes are lost (Duijf et al. 2013). As mitosis is an energy-demanding process and FTC-OV show an impaired ATP synthesis (Bonora et al. 2006, Savagner et al. 2001), it might be beneficial to lose larger chromosomes early in the progression process in order to save energy during mitosis. However, as we showed before in oncocytic adenocortical cancer and in oncocytic follicular thyroid cancer, disruptive mtDNA mutations, which might underlie a lack of ATP, do not solely explain the phenomenon of NHG (Corver et al. 2014). Disruptive mtDNA variations were also not found in the NHG cell lines FTC-133, -236 and -238. Together with the lack of a uniform nuclear gene variant in oncocytic tumors, the proliferation of mitochondria seen in both oncocytic follicular adenomas and carcinomas might partly be stress-related, possibly caused by a coupling defect (Savagner et al. 2001).

Hence, there seems to be a strong selection pressure on whole-chromosome losses after a pivotal asymmetric or reciprocal mitotic event in the development of FTC-OV, respectively. Other, probably random, whole-chromosome asymmetric mitotic events do not have a growth advantage or cells involved undergo apoptosis. Based on our observations, we propose a new progression model for (oncocytic) follicular thyroid tumors toward NHG with ROS and CHK2 activation as drivers of the process (Fig. 4C). The latter is characterized by a stepwise loss of whole chromosomes toward an NHG and cancer in which chromosome 22 might be a marker of progression.

Our data seem partly conflicting with earlier studies which demonstrated chromosomal copy number gains rather than losses in FTC-OV in most studies. Actually, previous findings are not that dissimilar from ours but were differently interpreted. We combined SNP array and multiparameter flow cytometric DNA content analyses, which provides allele-specific information and accurate copy numbers (Corver et al. 2008). The latter approach demonstrated that FTC-OV are dominated by whole-chromosome losses during tumor progression and that approximately 44% (Fig. 4A) of the cases in our cohort showed endoreduplication (genome doubling) of their entire NHG genome. For example, two copies of chromosomes
Table 3  A variety of genes mutated in FA-OV or FTC-OV cases (Hürthle cell carcinomas).

<table>
<thead>
<tr>
<th>Mutation/rearrangement fusions</th>
<th>Number of cases</th>
<th>Histology</th>
<th>Technique</th>
<th>% Positive</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>STK11*, TP53**</td>
<td>1</td>
<td>FTC-OV (with clear-cell features)</td>
<td>NGS (targeted)</td>
<td>100 (1/1)**</td>
<td>Wei et al. (2016)</td>
</tr>
<tr>
<td>KRAS*, HOOK3***, no fusions</td>
<td>6</td>
<td>FA-OV</td>
<td>RNA-seq</td>
<td>17 (1/6)*</td>
<td>Pagan et al. (2016)</td>
</tr>
<tr>
<td>BRAF</td>
<td>1</td>
<td>FTC-OV</td>
<td>PCR (BRAF)</td>
<td>100 (1/1)</td>
<td>Sinno et al. (2016)</td>
</tr>
<tr>
<td>PTTEN*, TP53**</td>
<td>12</td>
<td>FTC-OV</td>
<td>NGS (targeted)</td>
<td>25 (3/12)*</td>
<td>Wei et al. (2015)</td>
</tr>
<tr>
<td>TERT*, BRAF**</td>
<td>3</td>
<td>FTC-OV</td>
<td>PCR + direct sequencing</td>
<td>67 (1/3)*</td>
<td>Qasem et al. (2015)</td>
</tr>
<tr>
<td>MEN1* (EWSR1***, BRCA1***, MSH2***)</td>
<td>2</td>
<td>FTC-OV (fresh/ frozen)* shared in both tumors, **primary tumor, ***metastasis of the same tumor</td>
<td>Whole-genome sequencing Sanger sequencing for verification</td>
<td>100 (2/2)</td>
<td>Kasaian et al. (2015)</td>
</tr>
<tr>
<td>TSC2*, TP53*, FLCN*, mTOR**</td>
<td>1</td>
<td>ATC (from FTC-OV)</td>
<td>Whole-exome sequencing</td>
<td>100 (1/1)*</td>
<td>Wagle et al. (2014)</td>
</tr>
<tr>
<td>HRAS, NRAS, PAX8/ PPARγ*, RET/PTC</td>
<td>3</td>
<td>FTC-OV</td>
<td>RT-PCR</td>
<td>0 (0/12)</td>
<td>Mond et al. (2014)</td>
</tr>
<tr>
<td>BRAF, HRAS, KRAS, NRAS*</td>
<td>12</td>
<td>FA-OV</td>
<td>PCR</td>
<td>8.3 (1/12)*</td>
<td>Schulten et al. (2013)</td>
</tr>
<tr>
<td>TERT</td>
<td>17</td>
<td>FTC-OV</td>
<td>PCR</td>
<td>24 (4/17)</td>
<td>Landa et al. (2013)</td>
</tr>
<tr>
<td>BRAF, HRAS, KRAS, NRAS*, PIK3CA, PAX8/ PPARγ*, RET/PTC, NTRK1, AKAP9-BRAF rearrangement</td>
<td>8</td>
<td>FA-OV</td>
<td>Mass spectrometry genotyping (targeted)</td>
<td>0 (0/8), no rearrangements</td>
<td>0 (0/8)</td>
</tr>
<tr>
<td>PAX8/ PPARγ*, RET/PTC, NTRK1, AKAP9-BRAF rearrangement</td>
<td>19</td>
<td>FTC-OV</td>
<td>RT-PCR (rearrangements)</td>
<td>16 (3/19)*</td>
<td>de Vries et al. (2012)</td>
</tr>
<tr>
<td>TERT</td>
<td>3</td>
<td>FA-OV</td>
<td>FISH</td>
<td>33 (1/3)*</td>
<td>27 (4/17)**</td>
</tr>
<tr>
<td>EGRF, HRAS, KRAS, NRAS, PIK3CA*</td>
<td>11</td>
<td>FTC-OV</td>
<td>PCR (Sanger)</td>
<td>6.7 (1/15)*</td>
<td>Corver et al. (2012)</td>
</tr>
<tr>
<td>EGFR</td>
<td>1</td>
<td>FTC-OV (focally anaplastic)</td>
<td>PCR (followed by sequencing)</td>
<td>100 (1/1)</td>
<td>Hogan et al. (2009)</td>
</tr>
<tr>
<td>BRAF</td>
<td>2</td>
<td>FA-OV</td>
<td>PCR (followed by direct sequencing)</td>
<td>0 (0/21)</td>
<td>Musholt et al. (2008)</td>
</tr>
<tr>
<td>BRF, NRAS, PAX8/ PPARγ1</td>
<td>9</td>
<td>FTC-OV</td>
<td>PCR (followed by direct sequencing)</td>
<td>0 (0/9)</td>
<td>Di Cristofaro et al. (2006)</td>
</tr>
<tr>
<td>HRAS, KRAS*, NRAS</td>
<td>6</td>
<td>FTC-OV (familial)</td>
<td>PCR (followed by SSCP)</td>
<td>0 (0/6)</td>
<td>Tallini et al. (1999)</td>
</tr>
<tr>
<td>TSHR</td>
<td>4</td>
<td>FTC-OV</td>
<td>PCR (followed by SSCP)</td>
<td>25 (1/4)*</td>
<td>Spambalga et al. (1996)</td>
</tr>
</tbody>
</table>

A survey of the known literature on oncocytic adenoma and carcinoma was carried out. A plethora of techniques has been used and various mutations and/or rearrangements have been found by many different research groups. * or ** or ***, etc. corresponds to the percentage (x) and number of positive samples (y) of the total number of samples tested (z) in x(y/z) for a certain mutation/rearrangement/fusion in each study.

FISH, fluorescence in situ hybridization; NGS, next-generation sequencing; PCR, polymerase chain reaction; qPCR, quantitative polymerase chain reaction; RNA-seq, RNA sequencing; RT-PCR, reverse transcriptase polymerase chain reaction; SSCP, single-strand conformational polymorphism.
7 and 12 in a heterozygous state (AB) in a near-haploid (A) genomic background will be interpreted as gains by array-CGH. After endoreduplication also, the homozygous state of the residual chromosomes (AA) will not be seen by FISH.

Our observations regarding NHG in thyroid tumors have recently been confirmed (Kasaian et al. 2015) and have so far exclusively been found in NMTC with an oncocytic follicular histology or ATC that progressed from an FTC-OV (Wagle et al. 2014). However, the thyroid cancer cell lines FTC-133, -236 and -238 showed NHG but were reported to be derived from non-oncocytic FTC.

We now conclude that CHK2 activation via ROS is a dominant driver in the carcinogenesis of FTC-OV lesions with NHG, although we cannot rule out that the same or a comparable mechanism is active in the progression of other subsets of thyroid tumors.

Supplementary data
This is linked to the online version of the paper at https://doi.org/10.1530/ERC-17-0288.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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