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Accessibility
Paired Exome Analysis of Barrett’s Esophagus and Adenocarcinoma

Matthew D. Stachler1,2,*, Amaro Taylor-Weiner3,1, Shouyong Peng2, Aaron McKenna4, Agoston T. Agoston1, Robert D. Odze1, Jon M. Davison5, Katie S. Nason5, Massimo Loda1,2, Ignaty Leshchiner3, Chip Stewart3, Petar Stojanov3, Sara Seepo3, Michael S. Lawrence3, Daysha Ferrer-Torres6, Jules Lin6, Andrew C. Chang6, Stacey B. Gabriel3, Eric S. Lander3,7, David G. Beer6, Gad Getz3,8,#, Scott L. Carter3,9,#, and Adam J. Bass2,3,#

1Department of Pathology, Brigham & Women’s Hospital and Harvard Medical School, Boston, Massachusetts 02115, USA
2Department of Medical Oncology, Dana-Farber Cancer Institute, Boston, Massachusetts 02215, USA
3The Eli & Edythe L. Broad Institute, Cambridge, Massachusetts 02142, USA
4University of Washington, Seattle, Washington, 98104, USA
5University of Pittsburgh Cancer Institute, University of Pittsburgh, Pittsburgh, Pennsylvania, USA
6Section of Thoracic Surgery, University of Michigan, Ann Arbor, Michigan, USA
7Department of Biology, Massachusetts Institutes of Technology, Cambridge, Massachusetts, USA
8Department of Pathology, Massachusetts General Hospital, Boston, Massachusetts 02114, USA

Author Contributions:
MDS performed experiments and interpretation results. A.T.M., S.P., A.M., P.S., I.L., M.S.L., and S.L.C performed computational analysis. A.T.A. and R.D.O. performed pathologic slide review. J.M.D., K.S.N., D.F.T., J.L., A.C.C., and D.G.B. contributed samples and clinical annotation. M.L. contributed laser capture microdissection guidance and manuscript review. C.S., S.S., S.B.G and E.S.L organized and supervised sequencing. G.G. and S.L.C. and A.J.B. supervised all studies. M.D.S., A.T.M., S.L.C., G.G., and A.J.B. prepared the manuscript and all authors read and approved the final manuscript.

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Binary sequence alignment/map (BAM) files were deposited in the database of Genotypes and Phenotypes (phs000598).

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The authors declare no competing financial interests.

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Abstract

Barrett’s esophagus, is thought to progress to esophageal adenocarcinoma (EAC) through a step-wise progression with loss of CDKN2A followed by p53 inactivation and aneuploidy. Here, we present whole exome sequencing from 25 pairs of EAC and Barrett’s and five patients whose Barrett’s and tumor were extensively sampled. Our analysis revealed that oncogene amplification typically occurred as a late event and that TP53 mutations often occur early in Barrett’s progression, including in non-dysplastic epithelium. Reanalysis of additional EAC exome data revealed that the majority (62.5%) of EACs emerged following genome doubling and that tumors with genomic doubling had different patterns of genomic alterations with more frequent oncogenic amplifications and less frequent inactivation of tumor suppressors, including CDKN2A. These data suggest that many EACs emerge not through gradual accumulation of tumor suppressor alterations but rather through a more direct path whereby a TP53-mutant cell undergoes genome doubling, followed by acquisition of oncogenic amplifications.

Introduction

Barrett’s esophagus (BE), the intestinalization of the lower esophagus, develops in response to chronic gastric reflux and is the precursor to esophageal adenocarcinoma (EAC)\(^1,2,3\). While BE is estimated to exist in at least 1:100 adults\(^4\), relatively few progress to cancer. Those that do develop cancer are typically diagnosed at an advanced, incurable stage. Therefore, there is substantial interest in defining the molecular and genomic features of aggressive BE to enable means to prevent cancer or identify disease when most curable. The transformation of BE to EAC follows the progressive development of increasing grades of dysplasia, leading to invasive carcinoma. Several studies have characterized differences between BE, dysplasia, and EAC, looking at genomic copy-number\(^5–8\) and focused analysis of specific cancer-associated genes\(^5,9–12\). Through studies of BE, a linear model of progression has emerged whereby early non-dysplastic BE represents a clonal or polyclonal expansion, typically following inactivation of the tumor suppressor CDKN2A\(^5,7–10\). Further sub-clonal expansions are thought to occur, often leading to the emergence of a dysplastic clone with TP53 inactivation and other somatic alterations, including frequent genome doubling and increasing genomic disruption, leading to malignant transformation\(^7,9\). We sought to further clarify the process underlying transformation of BE to EAC by performing genomic analysis on BE and EAC samples derived from the same patient. We then extend this analysis to a cohort of previously sequenced EAC samples.

Results

Paired Barrett’s and esophageal adenocarcinoma analysis

We first performed whole exome sequencing (WES) on 25 patient-matched ‘trios’, including fresh-frozen EAC, BE, and non-malignant, distant gastric or esophageal squamous tissue as a germline comparator (Supplementary Table 1). All samples were obtained by
surgical resection from patients without prior chemo/radiotherapy, with BE intentionally isolated from a region not immediately adjacent to the tumor (when possible) during processing to avoid contamination of the BE with EAC cells. Upon pathologic review, 14 BE samples contained no dysplasia (BE\textsubscript{NoDys}) and 11 of the BE samples showed evidence of dysplasia (BE\textsubscript{Dys}). Of the 11 dysplastic samples, six contained changes consistent with high-grade dysplasia (HGD). Following somatic mutation calling (Supplementary File 1), we inferred the degree of shared ancestry of the paired BE and EACs based upon the number of shared mutations. In 11 of the 25 trios the specific region of sequenced BE appeared to be clonally unrelated to the sampled tumor as they lacked shared somatic mutations (Supplementary Fig. 1, Supplementary File 2). In addition, hierarchical clustering of the paired samples using somatic copy number alterations (SCNAs) failed to cluster these unrelated sample-pairs together (Supplementary Fig. 2). In the remaining 14 trios, the sampled regions of BE and EAC showed evidence of having emerged from a common neoplastic clone, as they shared 3.4 – 64% of coding point mutations with cancer cell fraction (CCF) of 1 (i.e., are present within all neoplastic cells in the tissue sample). Overall, we found no association between presence of dysplasia and whether the BE and EAC samples are clonally related (Fisher’s exact test, \(P=0.69\)). Among the six samples with HGD, five were clonally related to the EAC (\(P=0.18\)).

Our initial analyses compared the degree of genomic disruption between BE and EAC samples. The somatic mutation frequencies of BE ranged 1.3 – 5.4 mut/Mb – demonstrating that even non-dysplastic BE has mutation frequencies higher than many common invasive cancers (Fig. 1a). Mutational densities increased from BE\textsubscript{NoDys} (2.8 mut/Mb) to BE\textsubscript{Dys} (4.9 mut/Mb, Student t-test \(P=0.05\)), but were similar between BE\textsubscript{Dys} and EAC (4.1 mut/Mb, \(P=0.43\)). We also evaluated the base context of mutations in BE to query the presence of the common pattern of A to C transversions at the 3’ adenine at AA dinucleotides that we described in EAC\textsuperscript{13}. The predilection for these transversions was also present in BE (Fig. 1b), suggesting these mutations occur early during neoplastic progression, possibly caused by exposure to bile and acid reflux. Within the 14 clonally related cases, we separately analyzed the mutation spectra of the early shared mutations compared to those that were private to either the BE or the EAC. While there was a trend in increased A to C mutations private to BE, this was not statistically significant (Supplementary Fig. 3).

Our analysis of structural genomic changes revealed that, in contrast to point mutations, there was a greater increase in structural genomic disruption with progression to cancer. The mean number of focal deletions per sample increased steadily between BE\textsubscript{NoDys} (10.43), BE\textsubscript{Dys} (20.73) and the paired EAC samples (40.20, \(P<0.01\) for both) (Fig. 1c). More significantly, the number of amplifications was strongly associated with progression from BE\textsubscript{Dys} to EAC (Fig. 1d). BE\textsubscript{NoDys} and BE\textsubscript{Dys} averaged 0.42 and 0.91 amplifications per sample, respectively, while EACs averaged 8.44 amplifications per sample (\(P<0.001\) for both) suggesting that amplifications may be key mediators of oncogenic transformation. Even between BE with HDG and EAC, we still found a significant increase in amplifications, \(P=0.009\) (Supplementary Fig. 4). When we lowered amplification threshold to include lower-level gains the significant increase with progression persisted (Supplementary Fig. 5).
We evaluated specific oncogenic alterations within the trios, starting with tumor suppressor gene (TSG) alterations (Supplementary File 1, Supplementary Table 2). Within the 11 BE samples clonally unrelated to the paired EAC, only one harbored a TP53 mutation (Fig. 2a). While four of these BE cases possessed a homozygous CDKN2A deletion, the EACs that emerged in those patients lacked detectable somatic CDKN2A alterations. When we next evaluated the clonally related cases, we found that three of the four most distantly related trios (sharing 3.4%, 4.5%, and 7.8%, of mutations) had TP53 mutations shared between the BE and EAC (Fig. 2a, b), indicating that these TP53 mutations were among the earliest somatic mutations in the development of these tumors. All three of these tumors with early shared TP53 mutations were determined to have undergone whole genomic doubling (WGD) using the ABSOLUTE algorithm\cite{14}.

Furthermore, in these patients with evidence for an early TP53 mutation, shared CDKN2A somatic alterations were not observed. Seven of the 14 related BE/EAC cases had shared mutations in TP53 but not CDKN2A, which was either unaltered or had distinct inactivating events in the BE/EAC samples, suggesting that TP53 mutation was not preceded by CDKN2A inactivation in these cases (Fig. 2a, c). Only two of the 14 related cases appeared to clearly follow the classic model whereby the BE and EAC share a CDKN2A alteration but not a TP53 alteration, indicating that CDKN2A inactivation occurred before TP53. These results suggest that TP53 mutations may be an earlier event in BE pathogenesis in relation to other genomic alterations than previously recognized, often preceding (or occurring without) CDKN2A inactivation.

In contrast to the prevalent TSG alterations in BE, oncogenic activation events were far less prevalent in the sampled BE lesions, Fig. 3, even in those samples with advanced dysplasia or where the sampled BE appeared to be closely related to the cancer (Supplementary Fig. 6). High-level amplification of oncogenic cell-signaling proteins, cell-cycle modulators, and transcription factors were recurrently present in EACs but more infrequent in BE (Fig. 3, Supplementary Fig. 7 and 8). In addition, activating mutations of oncogenes were uncommon in BE, with only one known activating mutation identified, a PIK3CA E545Q mutation (Supplementary File 1), which was shared with the paired EAC. Oncogenic mutations in the 25 EAC were also uncommon, with two PIK3CA and one CTNNB1 hotspot mutation identified (Fig. 3). Together, oncogenes in cell signaling proteins (64% vs. 12%, \(P=0.0003\)), cell-cycle modulators (40% vs. 8%, \(P=0.0181\)), and transcription factors (64% vs. 20%, \(P=0.0037\)) were more likely to have an activating event in the EAC than the paired BE. These data suggest that while BE harbors common TSG inactivation; oncogene activation is a late step and may mediate transformation to cancer. In the patient sample (#63) available for confirmatory testing, fluorescent in-situ hybridization (FISH) for ERBB2 confirmed our sequencing analysis showing high level amplification in the EAC sample and no amplification within the BE tissue (Supplementary Fig. 8).

**Expanded multi-sample analysis via laser capture microdissection**

To better understand the progression of BE to EAC we identified 5 additional patients (Supplementary Table 3) with a broad field of BE who underwent surgical esophagectomies (without neoadjuvant therapy), from which we obtained paraffin-embedded samples.
representing multiple distinct stages of BE/EAC. We utilized laser capture microdissection to isolate normal tissue and pathologic samples spanning non-dysplastic BE, BE with low-grade dysplasia (LGD), BE with HGD, EAC, and nodal metastatic foci. The number of neoplastic samples characterized per patient ranged from 5 to 11 (Fig. 4 – 5). Laser capture microdissection allowed us to sample regions of BE in closer proximity to the tumor than feasible with the earlier trios from fresh tissue. With DNA from these newly dissected samples, we performed WES and processed data to identify somatic copy-number alterations, mutations, and genomic doubling (Supplementary Fig. 9). We then constructed phylogenetic trees indicating the evolutionary relationships between the subclones detected in each sample (Fig. 4 – 5; Supplementary Figs. 10 – 13). In three of the five cases all of the identified BE regions (both non-dysplastic and dysplastic) were clonally related to the EAC (Fig. 4). In the other two cases, all sampled regions of non-dysplastic BE were clonally unrelated to the EAC but we identified dysplastic BE samples related to the sampled tumor (Fig. 5). Most individually dissected tissue-samples were sufficiently diverged from one another such that no sharing of minor subclones occurred (mutations with CCF < 1 in two or more samples). However, patients 3 and 7 appeared to contain samples with partially overlapping subclones (Fig. 4f, Fig. 5b).

BE samples from two of the five esophagectomy cases supported our earlier observations that \(TP53\) mutations may occur earlier than previously thought in BE pathogenesis. In P1 and P7 (Fig. 4b, 4f) \(TP53\) missense mutations were shared by all sampled tissues, including regions of non-dysplastic BE (Supplementary Fig. 14). Furthermore, these two patients also demonstrated how oncogene amplification can be a late event mediating transformation. In P1, we identified a focal \(CDK6\) amplification present in the two EAC samples but not in the BE or HGD samples. However, in the \(TP53\)-mutant regions of non-dysplastic BE and HGD, we detected a \(GATA6\) amplification that is absent from the tumor (Fig. 4b) demonstrating that amplifications are not exclusive to EAC. In P7, we identified a high level \(KRAS\) amplification in the tumor and in a focus of HGD immediately adjacent to the cancer (HGD1; Fig. 4f). The \(KRAS\) amplification was notably absent from other regions of low and high-grade dysplasia more distant to the tumor (LGD1 and HGD2). FISH for \(KRAS\) confirmed these findings with high-level (>25 copies) amplification present in the EAC and absence of amplification in HGD2 or LGD1 (Supplementary Fig. 15).

Individually dissected samples within P7 also demonstrated substantial overlap of distinct subclonal populations. Sample LGD1 contained a major subpopulation (subclone 1a; CCF = 0.80 – 0.85) that was closely related to the last common ancestor of all neoplastic cells sampled in the patient. In addition, sample LGD1 contained a minor subpopulation (subclone 1b; CCF = 0.15 – 0.2) defined by 12 additional mutations, as well as single-copy loss of chromosomes 5q, 11p, and 13, and copy-neutral LOH of chromosomes 7 and 21 (Supplementary Fig. 9 – 12). Subclone 1c was descended from a cell closely resembling subclone 1b, with 10 additional mutations; subclone 1c was sampled only in HGD2. Subclone 1c closely resembled the common ancestor of all the subclones sampled in MET1, HGD1, EAC2, and EAC1 (none of which contained shared minor subclones with CCF<1). Sample BE1 contained two major subpopulations (subclone 2a; CCF = 0.4 – 0.6 and subclone 2b; CCF = 0.5 – 0.6). Subclone 2a was descended from a subclone closely
resembling subclone 1a, defined by 7 additional mutations. Subclone 2b was descended from a cell closely resembling subclone 2a, defined by 23 additional mutations (including \textit{AKT1} p.G10V) and gains of chromosomes 10 and 18p (Supplementary Fig. 9 – 12). We redrew the phylogenetic tree to represent the relationships between all subclones identified and overlaid the tissue-samples onto the tree (Fig. 4f).

P4 (Fig. 4c, 4d), exemplified a situation where a \textit{CDKN2A} deletion is present through all sampled regions of BE and EAC. We therefore inferred that a \textit{CDKN2A}-null progenitor gave rise to five distinct non-dysplastic BE subclones and one focus of LGD, all lacking \textit{TP53} alteration. However, we identified another subclonal branch in this case, consisting of five HGD/EAC samples all containing a \textit{TP53} mutation (p.R175H), focal low level (2 extra copies) amplification of \textit{MET}, and genomic doubling, consistent with a cancer in which \textit{TP53} mutation occurred later in clonal evolution, following \textit{CDKN2A} inactivation. Moreover, this case also supports the potential for oncogenic amplification to mediate transformation, as the one EAC sample harbored amplifications at \textit{CCNE1}, \textit{GATA6}, \textit{AKT2}, and 3q26 that were absent from all other samples. Samples HGD1, HGD2, and HGD3 all harbored a 7Mb gain with approximately 3 extra copies compared to a 4N baseline on 17q including \textit{ERBB2}, not present in the EAC.

Analysis of tissue from P3 and P6 (Fig. 5) demonstrated the potential for heterogeneity within the field of both BE and EAC. P3’s case contained three separate neoplastic regions, each clonally unrelated to the others. A first region contains an area of non-dysplastic BE with a focal \textit{CDKN2A} deletion. From this region emerged a segment of LGD with biallelic loss of \textit{ARID1A}. Separate from this region of BE were two clonally unrelated primary cancers (Supplementary Fig. 16). The sample EAC1 contained a mix of two distinct populations. Subclone 1 harbored 182 unique mutations (including \textit{SMARCA4}) and additional copy-number alterations, including a \textit{CCND1} amplification. EAC1 also contained a subpopulation of cells (subclone 2), that closely resembled the common ancestor of all subclones in the samples of more advanced disease, HGD1, EAC2, and Metastasis 1, with 125 mutations unique to their branch and unrelated to subclone 1. HGD1/EAC2 and Metastasis 1 also underwent WGD, which was not evident in EAC1/Subclone 1. Our focused analysis of this branch of advanced neoplastic tissues demonstrated how oncogene amplification can lead to transformation. The sample of HGD contained low-level segmental gains on chromosomes 6, 7, 8, and 17. The EAC and metastasis possessed additional amplifications, often occurring on top of these gains in the HGD. EAC2 harbored a more focal \textit{MYC} (chr 8) amplification and focal amplifications of 3q26 and \textit{MET} (chr 7) not identified in the region of HGD. In addition, Metastasis 1 (from a local lymph node) had an increased \textit{ERBB2} amplification (chr 17), demonstrating further heterogeneity.

Our analysis of patient 6 revealed one region of clonally distinct non-dysplastic BE as well as two EAC foci and two of HGD, which all emerged with WGD and a shared \textit{ATM} nonsense mutation. Interestingly, a focal \textit{CDKN2A} deletion was present in one of dysplastic samples (HGD1) but not in the HGD2 or EAC samples – another demonstration of a late \textit{CDKN2A} inactivation. The two EAC samples, both shared a \textit{MET} amplification not present in the dysplastic tissue.
Wider analysis of esophageal adenocarcinoma exomes

In order to validate our findings and better estimate the fraction of EAC cases that do not follow the conventional EAC evolution model, we reanalyzed whole-exome sequencing data from 144 microsatellite-stable EACs that we recently presented. Consistent with the conventional progression model, these tumors showed marked aneuploidy and harbored TP53 mutations in 97/144 (67%) of cases. However, we identified CDKN2A inactivation via inactivating mutations or homozygous deletions in only 39/144 (27%) of tumors.

We reanalyzed the EAC WES profiles using ABSOLUTE to make WGD calls and to determine whether specific mutations likely occurred prior to or following doubling. The majority (62.5%) of EACs showed evidence of WGD, corroborating previous findings. While WGD and non-WGD tumors showed different ploidy and genomic disruption, mutation densities were not different between these two groups (Supplementary Figs. 17, 18). TP53 mutations were distributed evenly across WGD (67% TP53 mutant) and non-WGD EACs (69% TP53 mutant). Within the WGD samples, 54/60 (90%) of the TP53 mutations were determined to have occurred prior to doubling. We compared the timing of TP53 mutations with that inferred for other mutated TSGs, including CDKN2A. TP53 was the only gene whose mutations were statistically more likely to have occurred prior to doubling (P=8.7×10⁻⁷; Supplementary Fig. 19, Supplementary Table 4) suggesting TP53’s role as an antecedent to genomic doubling.

Next, we searched for genomic characteristics that differentiate WGD and non-WGD EAC, finding early evidence for a different spectrum of TSP alterations. CDKN2A and SMAD4 mutations were found in only 8% and 1% of WGD EACs, respectively, but are present in 19% (P=0.065) and 20% (P=0.0001) of non-WGD tumors (Fig. 6a). Broadening our TSG analysis, we found TSG losses involving chromatin modification (P=0.005), cell cycle (P=0.027), and the TGF-β pathway (P=0.0001) were all more frequent in the non-GD EACs (Fig. 6b, Supplementary Tables 5 – 6). While we found that the mean number of deleted segments was not statistically different between WGD and non-WGD tumors, (29.13 vs. 30.93; P=0.28), there was a trend for increased homozygous deletions in the non-WGD samples (1.46 for non-WGD vs. 0.867 for WGD; P=0.104; Supplementary Fig. 20), as previously observed in serous ovarian cancer. The observation that tumors that emerge out of WGD have fewer somatic alterations of TSGs suggested that different alterations might drive transformation in these cases.

Therefore, we next evaluated patterns of oncogene activation, finding that alterations predicted to activate oncogenic signaling molecules, KRAS, RTKs, or PI3-kinase signaling, did not differ between the WGD and non-WGD groups (64% vs. 54%, P=0.22 for KRAS/RTKs and 9% vs. 13%, P=0.57 for PIK3CA) (Fig. 7). However, tumors with WGD showed more frequent amplification of oncogenic transcription factors (43 vs. 22%; P=0.012) and a trend towards higher rates of amplifications of cell-cycle mediators (40 vs. 24%; P=0.069). More frequent CCNE1 amplifications in genome doubled EACs (16% vs. 6%) paralleled recent analyses across tumor types which identified correlations of CCNE1 amplifications and WGD. These findings indicate that the absence or presence of WGD in BE modifies the most likely pathway available to undergo transformation to cancer.
Discussion

When the results of the EAC reanalysis is interpreted jointly with our earlier data on genomic analysis of paired BE/EAC cases, the results lead us to refine the previous model of the emergence of EAC from BE. EAC has been thought to follow the progressive accumulation of genomic alterations, starting with CDKN2A followed by expansion of a TP53 mutant dysplastic clone that is able to develop tetraploidy and genomic instability7,20. Consistent with this idea we confirm that BE harbors frequent TSG alterations, even in BE segments clonally unrelated to the cancer, corroborating reports of TSG inactivation in non-dysplastic BE21 and clonal diversity within BE22. Our data reinforce models that underscore the importance of TP53 mutation in neoplastic progression of BE23. However, our results also suggest that, in many cases, TP53 mutations occur earlier in the disease process relative to other alterations (including loss of CDKN2A) and can be detected in the non-dysplastic BE of those which progress to cancer. In addition, our data suggest that oncogene activation via amplification may often be a critical later event in transformation to invasive EAC. This sequence of events is in contrast to that seen in pancreatic or colorectal cancer where oncogene activation via mutation is believed to occur earlier, typically before TP53 inactivation24 – 27.

Our data therefore suggests that the traditional BE progression model conflates two general pathways for oncogenic transformation (Fig. 8). One pathway, starting similarly to the traditional model, appears to involve progressive accumulation of TSG losses (commonly CDKN2A and TP53 and also frequently including SMAD4 and alterations of chromatin modifying enzymes), leading to genomic instability and oncogenic amplifications without an antecedent WGD. Despite the prevalence of clonal expansions of BE tissues with somatic inactivation of TSGs such as CDKN2A and ARID1A, a minority of EACs appear to emerge following such a path without WGD. Instead, the majority of EACs apparently develop following expansion of a TP53 mutant clone that undergoes WGD, with WGD predominately seen in tissues with dysplasia. Genomic doubling has been documented to facilitate the acquisition of genomic instability16,28,29. Consistent with this, those tumors that emerged following WGD harbored marked genomic disruption and oncogene amplification, with amplifications of known oncogenes observed in 77/90 WGD EAC samples. Following WGD, homozygous inactivation of tumor suppressors becomes more difficult due the additional number of events required14. Once WGD occurs, the more expedient pathway to transformation is thus likely that of acquisition of oncogene activation via structural genomic instability. The predilection for instability following WGD16,29 – 31 likely contributes to catastrophic genomic disruptions resulting in a large number of copy number alterations, as recently identified by whole genome sequencing of EAC32. This alternative pathway to transformation whereby a WGD dysplastic clone acquires oncogene activation via amplification is supported by our findings that oncogene amplification is typically a later event in the progression to EAC. Such episodes of genomic disruption could lead to positive selection for distinct amplifications that bypass the need for the loss of tumor-suppressors, as more frequently occurs in non-WGD EACs (Fig. 3). We note that it is also possible that WGD cells are better able to survive catastrophic genomic disruption (e.g. since such events are unlikely to generate complete gene knockouts), and may be less
subject to negative selection. While our limited dataset did not identify differences in clinical stage between WGD and non-WGD EACs (Supplementary Fig. 21), future studies will be needed to identify whether tumors following these distinct paths have other distinguishing clinical features.

Both WGD and TP53 mutations have been recognized for over a decade to be risk-factors for the development of EAC in patients with BE\(^7,23,33,34\). Here we are able to refine the conventional model of how these tumors emerge, finding that TP53 mutations may be earlier events than previously recognized. Following TP53 mutation many EACs may follow a distinct pathway to cancer involving WGD with subsequent transformation to cancer via catastrophic aneuploidy and oncogene amplification. Our model is therefore consistent with a recent complementary report which characterized the copy-number profiles of serially collected BE biopsies by Li et al. in which the authors noted that in the 24 months prior to cancer diagnosis a marked increase in DNA content occurred, suggestive of WGD\(^7\).

Our refined model positing a potentially more rapid path to transformation following acquisition of a TP53 mutation and WGD may help explain the failure of endoscopic screening of BE patients to prevent cancer diagnoses and deaths. Screening strategies are largely premised upon the concept that BE is at risk of gradual accumulation of genomic alterations leading to progression to cancer. This model would predict that just as aging predisposes to cancer, duration of BE would also enhance cancer risk. However, contrary to this concept, studies of population cohorts of patients with diagnoses of BE show that the majority of EACs are detected within the first two to three years of initial endoscopic diagnosis of BE, even when the incident endoscopy fails to identify dysplasia or cancer\(^35–38\).

If most EACs emerge from BE following a more rapid process involving TP53 mutation and emergence of an unstable genomically doubled intermediate, new diagnostic strategies may be required that seek out these TP53-mutant precursors and intermediates as a means of detecting and preventing this deadly disease.

**Online Methods**

**Sample selection and DNA extraction (fresh frozen)**

Samples were obtained with documented informed consent and institutional IRB approval. Fresh frozen samples were obtained at the time of surgical esophagectomy from patients with a diagnosis of esophageal adenocarcinoma and without neoadjuvant therapy from the University of Michigan. Samples of BE were selected not immediately adjacent to the tumor (when possible) to minimize tumor contamination within the BE sample. Hematoxylin and eosin (HE) stained slides from 27 cases were examined by a pathologist with sub-specialty training in gastro-intestinal pathology. Slides from BE tissue were classified as either non-dysplastic or dysplastic. Samples with high and low grade dysplasia were combined for primary analysis due to overall low numbers and the diagnostic challenges and controversies associated with distinguishing between these entities (especially on frozen section slides). Additional subanalyses looked separately at these diagnoses. Two cases were excluded from further analysis as one case contained too low of a percentage of BE and one BE tissue was contaminated with invasive tumor. DNA was extracted using phenol chloroform and ETOH precipitation and quantified using Picogreen dsDNA Quantification Reagent (Invitrogen).
Sample selection and DNA extraction (formalin fixed and paraffin embedded)

With documented informed consent and institutional IRB approval, formalin fixed paraffin embedded (FFPE) esophagectomy samples without neoadjuvant therapy were identified in the pathology archives of the University of Pittsburgh Medical Center and Brigham and Women’s Hospital. HE slides were reviewed by two gastrointestinal pathologists to determine consensus areas of BE, BE with low grade dysplasia, BE with high grade dysplasia, and esophageal adenocarcinoma. If uncertainty for a diagnosis was present, a third pathologist reviewed the sample. Any sample without a consensus diagnosis was eliminated from analysis. Ten 8 micron sections were cut onto PEN membrane frame slides (Life Technologies, Grand Island, NY) bracketed by standard slides for HE staining. The frame slides were stained using Arcturus paradise plus stain (Life Technologies) following the manufacturers recommendations. The areas of interest were microdissected using the ArcturusXT laser capture microdissection Instrument (Life Technologies). DNA was isolated using the Qiagen (Valencia, CA) FFPE DNA isolation kit following the manufacturers protocol with the exception that the tissue was digested with proteinase K overnight. DNA was quantified using Picogreen dsDNA Quantification Reagent.

Whole exome sequencing

Whole-exome capture libraries were constructed from 100ng of DNA following shearing, end repair, phosphorylation and ligation to barcoded sequencing adapters. DNA was size-selected for lengths between 200–350bp and subjected to exonic hybrid capture using SureSelect v2 Exome bait (Agilent). Samples were multiplexed and sequenced on multiple Illumina HiSeq flow cells. Mean target exome coverage of 95x was achieved in the Barrett’s DNA, 85x in the neoplastic DNA, and 87x in the normal tissue.

Sequencing data processing

Exome sequence data processing and analysis were performed using Broad Institute pipelines as previously described. A BAM file aligned to the hg19 human genome build was generated from sequencing reads for each sample by the “Picard” pipeline.

Mutation calling

The MuTect algorithm was used to identify somatic mutations. We required a minimum of 14 reads covering a site in the tumor and 8 in the normal for declaring a site is adequately covered for mutation calling. We determined the lowest allelic fraction at which somatic mutations could be detected on a per-sample basis, using estimates of cross-contamination from the ContEst pipeline. Small somatic insertions and deletions were detected using the Indelocator algorithm after local realignment of tumor and normal sequences. All somatic mutations detected by WES were analyzed for potential false-positive calls by performing a comparison to mutation calls from a panel of 2,500 germline DNA samples. Mutations found in 2% of the germline samples or 2% of sequencing reads were removed from analysis.

Because of our goal of comparing the presence and absence of mutations between distinct samples taken from the same patient, we utilized a tool designed to specifically query evidence for mutations or insertions/deletions called in one sample for evidence of their
presence, even at low allelic fraction, in other samples from the same patient. The strong prior of having been called de-novo in one sample allows for more sensitive detection in other related samples. This method, termed ‘force calling’, utilizes outputs from MuTect and Indelocator to generate an aggregate set of somatic events for each patient. Then, it adopts Samtools to count the number of reads supporting the reference or alternate alleles at those sites in the other matched samples. Reads are considered if they are from unique pairs, have a base quality at the site of interest of greater than or equal to 20, and read quality greater than or equal to 5.

Mutation annotation

Somatic single-nucleotide variants, insertions, and deletions were annotated using Oncotator (www.oncotator.org), which uses information from publicly available databases.46–50.

Calculation of total and allelic copy-numbers from whole exome sequencing data

Genome-wide copy-ratio profiles were inferred using CAPSEG. Read-depth at informative capture targets in tumor samples was calibrated to estimate copy-ratio using depths observed in a panel of normal (non-cancer) diploid genomes. The resulting copy-ratio profiles were then segmented using the circular binary segmentation (CBS) algorithm. Allelic copy-number analysis was then performed by examination of alternate and reference read counts at heterozygous SNP positions (as determined by analysis of the matched normal sample). These counts were used to infer the contribution of the two homologous chromosomes to the observed copy-ratio in each segment. Further analysis of change-points in these allelic-ratios was performed using PSCBS, refining the segmentation. Finally, for each segment, we combined the copy-ratio and allelic data to derive allelic copy-ratios, which were input for analysis with ABSOLUTE.14

The ABSOLUTE computational tool (v1.2) was used to provide computational estimates of several parameters for each neoplastic sample in this study. These estimates include: (i) the purity of each sample (fraction of nuclei in the sample originating from tumor or BE); (ii) the average ploidy of the cancer or BE genome; (iii) the presence of antecedent genomic doubling for each genome; and (iv) the absolute allelic copy-number across the genome. ABSOLUTE takes as input the segmented allelic copy-number ratio data (as described above) as well as the allele fractions of somatic point mutations (aberrant reads as a ratio of total reads covering the locus), and then determines possible combinations of tumor purity, ploidy and antecedent genomic doubling which fit the allelic copy-number ratio data and point-mutation variant allele fraction (VAF; Supplementary Fig. 9). The ABSOLUTE solutions were reviewed manually to maximize concordance with the data (A.T.W.).

Calculation of point-mutation CCF distributions

For each somatic mutation, we computationally estimated the fraction of neoplastic cells within a specific DNA sample that harbors the mutation, i.e. its cancer cell fraction (CCF). This fraction is represented as a distribution between 0 and 1.14,53,54 A CCF value of 1 corresponds to a mutation present in 100% of neoplastic cells in a sample. A CCF value of <1 indicates that the mutation is present in a subset of the neoplastic cells in a sample and

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thus is subclonal. Probability distributions for CCF values were computed by correcting mutant and reference read counts from Illumina sequencing for the estimated sample purity and local copy-number\textsuperscript{14} (Supplementary Fig. 9), as previously described.\textsuperscript{53,54}

After the initial determination of a CCF distribution for each mutation, we then perform an additional analysis to refine our CCF estimates, based on the assumption that each neoplastic sample contained a small (but unknown) number of distinct populations defined by mutations that share the same CCF. This clustering is performed using Bayesian clustering that jointly estimates the CCF values and number of populations based upon the set of CCF distributions from each sample. This technique utilizes sampling from a mixture of Dirichlet processes using a Monte Carlo Markov Chain (MCMC) sampler, as previously described (Supplementary Fig. 11a, b)\textsuperscript{53,54}. We used 250 MCMC iterations where the 125 initial were discarded as ‘burn-in’. A prior over the number of mutation clusters in a given sample was specified using a negative binomial distribution ($r=10$, $\mu=3$), which favored 1–5 clusters (Supplementary Fig. 11d). A partition of mutations was obtained based on how often each pair of mutations was assigned to the same cluster during the MCMC simulation (after convergence). A distance metric was generated from the inverse of each pair-count, and hierarchical clustering was performed using complete linkage. The resulting tree was divided into $k$ clusters, with $k$ chosen as the lowest number of sampled clusters in the MCMC (Supplementary Fig. 11c, d).

**Calculation of statistical power for detection of shared somatic mutations**

To calculate power, we considered the expected variant allele fraction (VAF) of a point mutation with CCF=1 and multiplicity=1, given the sample purity and local copy number. For mutations detected in only a subset of the tissue samples comprising a given case, we calculated the *paired*-detection power. Because shared mutations with a single supporting-read matching the called allele were called by the forced-calling procedure (described above), we calculated power as the probability of observing one or more such read, given its expected VAF and sequence coverage.

**Relatedness calculations**

Relatedness was calculated by taking the total number of shared mutations that were present at CCF=1 in both samples divided by the sum of the total number of with CCF=1 mutations in either sample (Supplementary Fig. 1). The relatedness using all mutations (CCF=1 and CCF<1) was also calculated and reported in Supplementary Table 1. We also searched for evidence of shared copy-number aberrations, which would provide independent support for a potential common origin in a EAC/BE pair beyond shared point mutations. For each segment in the EAC samples with copy number greater than 2.5 or less than 1.5 we looked at the matched Barrett’s sample to see if there is also a segment greater than 2.5 or less than 1.5 that is 50% mutually overlapping with the segment in the EAC. If so, it was called shared. Using this analysis, we did not find any additional shared EAC/BE pairs not detected based upon shared somatic mutations.
Phylogenetic inference

Tumors can exhibit genetic heterogeneity both across different regions\textsuperscript{55–58} and within single cancer-tissue samples.\textsuperscript{14,53,54,59,60,61} Heterogeneity within individual tissue samples presents a difficulty to standard phylogenetic inference algorithms, which typically distinguish only between the presence or absence of mutations in each sample. These algorithms attempt to construct phylogenetic trees relating each tissue sample, which may not accurately reflect the evolutionary relationships between the neoplastic cell-populations represented in the tissue samples. For example, shared mutations (present in multiple samples) due to overlapping subclonal populations would be mistaken as evidence for shared ancestry between the samples.

To address this difficulty, we utilized quantitative information about each mutation’s prevalence in each neoplastic tissue sample (CCF) in order to determine whether the tissue samples were sufficiently diverged from one another such that no detectable overlap of minor subclones (CCF < 1) occurred, a scenario we term the branched-sibling model. In this scenario, it is valid to construct standard phylogenetic trees relating each tissue sample, with minor subclones (CCF < 1) private to each tissue sample represented as subtrees (“microphylogenies”) grafted on to each sample tip. The branched-sibling scenario implies that such trees accurately represent the evolutionary relationship of all subclonal populations detected in the sampled neoplastic tissues. A corollary of the branched-sibling model is that all mutations shared in two or more samples must have CCF=1 wherever they are present (Supplementary Fig. 12a, b). Thus, the appearance of mutations shared in two or more samples with CCF<1 in any of them either represents technical artifact or constitutes evidence that the branched-sibling approximation is not an accurate description of those samples.

We constructed phylogenetic trees representing the evolutionary relationship between the neoplastic tissue-samples sequenced from each patient using a semiautomated four-stage process, described below. First, we searched for the optimal tree that would explain the observed matrix of binary point-mutation presence or absence data in each sample, given the standard phylogenetic assumptions that specific mutations arise uniquely in each patient and that there were negligible rates of mutation loss (e.g. due to chromosomal deletion of a mutant allele). We searched for the phylogenetic tree with maximum parsimony using the standard parsimony-ratchet method.\textsuperscript{52}

Second, we applied the Bayesian CCF-clustering procedure described above to each sample individually, retaining all mutations provisionally called with > 0 supporting reads in that sample. A single pseudo-count observation was added corresponding to a cluster at CCF=1. We then identified all provisional mutation calls (> 0 supporting reads) made in at least two samples of the case that were assigned to a CCF cluster with posterior mode < 1.0 (Supplementary Fig. 12). These mutation calls represent either sequencing artifacts or evidence for overlapping minor subclones in the sampled neoplastic tissues (violating the branched-sibling approximation). We rejected such sites if the number of supporting reads was < 3, and this modified matrix of mutation calls was then used to assign each mutation to a branch of the phylogenetic tree (Supplementary Fig. 12b).
We also distinguished mutations that were underpowered for detection in some samples (as described above). For each sample, the number of mutations in each category is shown in Supplementary Fig. 12c. Assignment of gene-level SCNAs to branches was performed in a similar manner (Supplementary Fig. 12d).

Third, we refined the tips of each phylogenetic tree by distinguishing between private mutations that occurred in all neoplastic cells of given sample (CCF=1) compared to those that occurred in only a minor subclone (CCF < 1) specific to that sample. For this distinction we applied Bayesian clustering techniques (described above) to the mutations identified only in that sample. We added \( N \) pseudo-count observations of CCF=1, with \( N \) representing the number of mutations called in >1 samples of the case that were also called in the sample being considered. This process partitioned the private mutations in each sample into putative subclones with common CCF values (Supplementary Fig. 10). We modified the phylogenetic trees by replacing each (non-germline) tip with the subtree consistent with the maximally branching microphylogeny respecting the rule that the sum of sibling-subclone CCF values cannot exceed that of their most recent common ancestor; Fig. 4 and 5, Supplementary Fig. 13).

Fourth, we examined whether evidence that the branched-sibling model was not an adequate approximation of the sampled neoplastic tissues could be discerned. For each case, we analyzed the two-dimensional CCF distributions of point mutations for all unique tissue-sample pairs (Supplementary Fig. 11) using a 2D version of the Bayesian clustering algorithm described above.\(^63\) Examination of these data revealed robust clusters of mutations with CCF=1 in both samples of clonally related pairs. In addition, many pairs harbored mutations with CCF=1 in one sample that were undetected in the paired sample (and vice-versa). Most samples also harbored mutations with CCF<1 that were undetected in the paired sample. Furthermore, for most patients only a small fraction of mutations with CCF<1 appeared to be detected in both sample-pairs, and did not tend to form strong clusters (consistent with sequencing artifacts; Supplementary Fig. 11). Taken together, these observations implied that most tissue samples were well approximated by the branched-sibling model (Supplementary Fig. 11), since true overlap of minor populations from distinct subclonal branches would tend to displace the CCF values of mutations private to each sample, so that none would have CCF=1.

In the two cases where evidence contradicting the branched-sibling model was observed, phylogenetic trees were manually adjusted (as described below) to accurately reflect the evolutionary relationship between the different clonal lineages as shown in Fig. 4, 5. This was done in a manner analogous to that described in a recent report\(^61\); here we extended similar logic to the scenario where the same subclone was present in multiple tissue samples. Detailed analysis of mutation CCFs for each patient, including the automatically generated phylogenetic trees (prior to manual adjustment), are available in Supplementary Files 3, 4.

For patients 4 and 6, nearly every shared mutation had CCF=1. For patient 1, the shared mutations with CCF<1 did not appear to form a strong cluster, or to displace other clusters away from CCF=1. We therefore assumed that these mutations did not constitute strong
evidence of a branched-sibling violation, and accepted the automatically produced phylogenies for these cases.

For patient 3, we detected a minor population in sample EAC1 (subclone1; CCF=0.2) defined by 45–200 mutations (which were present at CCF=1 in the shared MET1/HGD1/EAC2 branch. We adjusted the phylogenetic tree (Fig. 5b) to move EAC1 onto a distinct branch from these samples in order to represent the dominant subclone (subclone2; CCF=0.8) in EAC1, which had an evolutionary origin distinct from that of subclone1. In addition, samples HGD1 and MET1 appeared to share a small subclone (CCF=0.05) defined by 5 mutations that did not appear elsewhere in the case. We removed the shared branch defined by these mutations from the phylogeny as they did not reflect shared recent ancestry of the dominant subclones in HGD1 and MET1 (Fig. 5b). For patient 7, the details regarding the phylogenetic tree generation are detailed in the main text.

**Fluorescent in-situ hybridization**

In P3, we wanted to confirm the genome doubling status in two apparent genomically unrelated tumors. ABSOLUTE predicted that one of the samples has undergone a genomic doubling event, whereas the other did not. We performed FISH analysis co-labeling a 4 μm FFPE section with two differently labeled centromeric markers for chromosomes that are relatively stable in EAC (CEP 2 and CEP 4). After review and identification of the areas of interest, the probes were enumerated by two senior research technologists within the Brigham and Women’s cytogenomics core facility. Each technologist counted 50 cells per sample and the number of CEP 2 and CEP 4 probes per cell were averaged. In P7, we performed FISH for KRAS and CEP 12 from samples LGD1, HGD2, and EAC1 to confirm the finding that the amplification was present EAC1 but not the premalignant lesions. FISH was performed on the paired samples BE 63 and EAC 63 for ERBB2 and CEP 17. All FISH studies were performed with standard techniques. The gene to CEP ratio was calculated and a ratio of greater than 2 was considered positive.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**

1. Nehra D, Howell P, Williams CP, Pye JK, Beynon J. Toxic bile acids in gastro-oesophageal reflux disease: influence of gastric acidity. Gut. 1999; 44:598–602. [PubMed: 10205192]
2. Wild CP, Hardie LJ. Reflux, Barrett’s oesophagus and adenocarcinoma: burning questions. Nat Rev Cancer. 2003; 3:676–84. [PubMed: 12951586]

3. Lagergren J, Bergstrom R, Lingren A, Nyren O. Symptomatic Gastrosophageal Reflux as a Risk Factor for Esophageal Adenocarcinoma. NEJM. 1999; 340:825–831. [PubMed: 10080844]

4. Ormsby AH, et al. The Location and Frequency of Intestinal Metaplasia at the Esophagogastric Junction in 223 Consecutive Autopsies: Implications for Patient Treatment and Preventive Strategies in Barrett’s Esophagus. Mod Pathol. 2000; 6:614–620. [PubMed: 10874664]

5. Galipeau PC, Prevo LJ, Sanchez Ca, Longton GM, Reid BJ. Clonal Expansion and Loss of Heterozygosity at Chromosomes 9p and 17p in Premalignant Esophageal (Barrett’s) Tissue. JNCI J Natl Cancer Inst. 1999; 91:2087–2095. [PubMed: 10601379]

6. Gu J, et al. Genome-wide catalogue of chromosomal aberrations in barrett’s esophagus and esophageal adenocarcinoma: a high-density single nucleotide polymorphism array analysis. Cancer Prev Res. 2010; 3:1176–1186.

7. Li X, et al. Temporal and spatial evolution of somatic chromosomal alterations: A case-cohort study of Barrett’s esophagus. Cancer Prev Res. 2013

8. Li X, et al. Single nucleotide polymorphism-based genome-wide chromosome copy change, loss of heterozygosity, and aneuploidy in Barrett’s esophagus neoplastic progression. Cancer Prev Res. 2008; 1:413–423.

9. Reid BJ, et al. Barrett’s esophagus: ordering the events that lead to cancer. Eur J cancer Prev. 1996; 5:57–65. [PubMed: 9061296]

10. Wong DJ, et al. p16 INK4a Lesions Are Common, Early Abnormalities that Undergo Clonal Expansion in Barrett’s Metaplastic Epithelium p16 INK4a Lesions Are Common, Early Abnormalities that Undergo Clonal Expansion. Cancer Res. 2001; 61:8284–8289. [PubMed: 11719461]

11. Zhang S, Wang XI. SIRT1 is a Useful Biomarker for High-Grade Dysplasia and Carcinoma in Barrett’s Esophagus. 2013; 43:373–377.

12. Paulson TG, et al. p16 mutation spectrum in the premalignant condition Barrett’s esophagus. PLoS One. 2008; 3:e3809. [PubMed: 19043591]

13. Dulak AM, et al. Exome and whole genome sequencing of esophageal adenocarcinoma identifies recurrent driver events and mutational complexity. Nat Genet. 2013; 45:1–21. [PubMed: 23268125]

14. Carter SL, et al. Absolute quantification of somatic DNA alterations in human cancer. Nat Biotechnol. 2012; 30:413–21. [PubMed: 22544022]

15. Dulak AM, et al. Gastrointestinal adenocarcinomas of the esophagus, stomach, and colon exhibit distinct patterns of genome instability and oncogenesis. Cancer Res. 2012; 72:4383–93. [PubMed: 22751462]

16. Fujiwara T, et al. Cytokinesis failure generating tetraploids promotes tumorigenesis in p53-null cells. Nature. 2005; 437:1043–7. [PubMed: 16222300]

17. Davoli T, de Lange T. Telomere-driven tetraploidization occurs in human cells undergoing crisis and promotes transformation of mouse cells. Cancer Cell. 2012; 21:765–76. [PubMed: 22698402]

18. Zack TI, et al. Pan-cancer patterns of somatic copy number alteration. Nat Genet. 2013; 45:1134–1140. [PubMed: 24071852]

19. Etemadmoghadam D, et al. Resistance to CDK2 inhibitors is associated with selection of polyploid cells in CCNE1-amplified ovarian cancer. Clin Cancer Res. 2013; 19:5960–71. [PubMed: 24004674]

20. Maley CC, et al. Selectively advantageous mutations and hitchhikers in neoplasms: p16 lesions are selected in Barrett’s esophagus. Cancer Res. 2004; 64:3414–3427. [PubMed: 15150093]

21. Weaver MJ, et al. Ordering of mutations in preinvasive disease stages of esophageal carcinogenesis. Nat Genet. 2014; 8:837–43. [PubMed: 24952744]

22. Maley CC, et al. Genetic clonal diversity predicts progression to esophageal adenocarcinoma. Nat Genet. 2006; 38:468–73. [PubMed: 16565718]

23. Reid BJ, et al. Predictors of Progression in Barrett’s Esophagus II: Baseline 17p (p53) Loss of Heterozygosity Identifies a Patient Subset at Increased Risk for Neoplastic Progression. Am J Gastroenterol. 2001; 96:2839–2848. [PubMed: 11693316]
24. Van Wyk R, et al. Somatic mutations of the APC, KRAS, and TP53 genes in nonpolypoid colorectal adenomas. Genes Chromosomes Cancer. 2000; 27:202–8. [PubMed: 10612810]

25. Prestlow T, Prestlow T. No Mutant KRAS in aberrant crypt foci (ACF): Initiation of colorectal cancer? Biochim Biophys Acta. 2005; 1756:83–96. [PubMed: 16219426]

26. Lüttges J, et al. The K-ras mutation pattern in pancreatic ductal adenocarcinoma usually is identical to that in associated normal, hyperplastic, and metaplastic ductal epithelium. Cancer. 1999; 85:1703–10. [PubMed: 1022563]

27. Deramaudt T, Rustgi A. Mutant KRAS in the initiation of pancreatic cancer. Biochim Biophys Acta. 2005; 1756:97–101. [PubMed: 16169155]

28. Gordon DJ, Resio B, Pellman D. Causes and consequences of aneuploidy in cancer. Nat Rev Genet. 2012; 13:189–203. [PubMed: 22269907]

29. Dewhurst SM, et al. Tolerance of whole-genome doubling propagates chromosomal instability and accelerates cancer genome evolution. Cancer Discov. 2014; 4:175–85. [PubMed: 24436049]

30. Davoli T, de Lange T. The causes and consequences of polyploidy in normal development and cancer. Annu Rev Cell Dev Biol. 2011; 27:585–610. [PubMed: 21801013]

31. Ganem N, et al. Cytokinesis failure triggers hippo tumor suppressor pathway activation. Cell. 2014; 158:833–48. [PubMed: 25126788]

32. Nones K, et al. Genomic catastrophes frequently arise in esophageal adenocarcinoma and drive tumorigenesis. Nat Commun. 2014; 5:5224. [PubMed: 25351503]

33. Rabinovitch P, Reid B, Haggit R, Norwood T, Rubin C. Progression to cancer in Barrett’s esophagus is associated with genomic instability. Lab Invest. 1989; 60:65–71. [PubMed: 2911184]

34. Davelaar AL, et al. Ablation of TP53 detected by combining immunohistochemistry and DNA-FISH improves Barrett’s esophagus progression prediction: A prospective follow-up study. Genes Chromosomes Cancer. 2015; 54:82–90. [PubMed: 25284618]

35. Bytzer P, Christensen PB, Damkier P, Vinding K, Seersholm N. Adenocarcinoma of the Esophagus and Barrett’s Esophagus: A Population-Based Study. 1999; 94:1–6.

36. Westona P, et al. Long-term follow-up of Barrett’s high-grade dysplasia. Am J Gastroenterol. 2000; 95:1888–93. [PubMed: 10950031]

37. Corley D, Levin TR, Habel L, Weiss NS, Buffler P. Surveillance and survival in Barrett’s adenocarcinomas: A population-based study. Gastroenterology. 2002; 122:633–640. [PubMed: 11874995]

38. Hvid-Jenson F, Pedersen L, Drewes A, Sorensen H, Funch-Jensen P. Incidence of adenocarcinoma among patients with Barrett’s esophagus. N Engl J Med. 2011; 365:1375–1383. [PubMed: 21995385]

39. Fisher S, et al. A scalable, fully automated process for construction of sequence-ready human exome targeted capture libraries. Genome Biol. 2011; 12:R1. [PubMed: 21205303]

40. Berger MF, et al. The genomic complexity of primary human prostate cancer. Nature. 2011; 470:214–20. [PubMed: 21307934]

41. Chapman M, Lawrence M, Keats J. Initial genome sequencing and analysis of multiple myeloma. Nature. 2011; 471:467–472. [PubMed: 21430775]

42. Barbieri CE, et al. Exome sequencing identifies recurrent SPOP, FOXA1 and MED12 mutations in prostate cancer. Nat Genet. 2012; 44:685–9. [PubMed: 22610119]

43. Transky N, et al. The mutational landscape of head and neck squamous cell carcinoma. Science. 2011; 333:1157–60. [PubMed: 21798893]

44. Cancer genome action network. Integrated genomic analyses of ovarian carcinoma. Nature. 2011; 474:609–15. [PubMed: 21720365]

45. Cibulskis K, et al. ContEst: estimating cross-contamination of human samples in next-generation sequencing data. Bioinformatics. 2011; 27:2601–2. [PubMed: 21803805]

46. Fujita, Pa, et al. The UCSC Genome Browser database: update 2011. Nucleic Acids Res. 2011; 39:D876–82. [PubMed: 20959295]

47. Sherry ST, et al. dbSNP: the NCBI database of genetic variation. Nucleic Acids Res. 2001; 29:308–11. [PubMed: 11125122]
48. Griffith OL, et al. ORegAnno: an open-access community-driven resource for regulatory annotation. Nucleic Acids Res. 2008; 36:D107–13. [PubMed: 18006570]
49. UniProt Consortium. Ongoing and future developments at the Universal Protein Resource. Nucleic Acids Res. 2011; 39:D214–9. [PubMed: 21051339]
50. Forbes, Sa, et al. COSMIC: mining complete cancer genomes in the Catalogue of Somatic Mutations in Cancer. Nucleic Acids Res. 2011; 39:D945–50. [PubMed: 20952405]
51. Venkatraman ES, Olshen AB. A faster circular binary segmentation algorithm for the analysis of array CGH data. Bioinformatics. 2007; 23:657–63. [PubMed: 17234643]
52. Olshen AB, et al. Parent-specific copy number in paired tumor-normal studies using circular binary segmentation. Bioinformatics. 2011; 27:2038–46. [PubMed: 21666266]
53. Landau DA, et al. Evolution and impact of subclonal mutations in chronic lymphocytic leukemia. Cell. 2013; 152:714–726. [PubMed: 23415222]
54. Lohr JG, Stojanov P, Carter SL, Cruz-gordillo P, Lawrence MS. Widespread Genetic Heterogeneity in Multiple Myeloma: Implications for Targeted Therapy. Cancer Cell. 2014; 25:91–101. [PubMed: 24434212]
55. Campbell PJ, et al. The patterns and dynamics of genomic instability in metastatic pancreatic cancer. Nature. 2010; 467:1109–1113. [PubMed: 20981101]
56. Gerlinger M, et al. Intratumor Heterogeneity and Branched Evolution Revealed by Multiregion Sequencing. N Engl J Med. 2012; 366:883–892. [PubMed: 22397650]
57. Liu W, et al. Copy Number Analysis Indicates Monoclonal Origin of Lethal Metastatic Prostate Cancer. Nat Med. 2009; 15:559–565. [PubMed: 19363497]
58. Navin N, et al. Tumour evolution inferred by single-cell sequencing. Nature. 2011; 472:90–94. [PubMed: 21399628]
59. Shah SP, et al. Mutational evolution in a lobular breast tumour profiled at single nucleotide resolution. Nature. 2009; 461:809–813. [PubMed: 19812674]
60. Nik-Zainal S, et al. Mutational processes molding the genomes of 21 breast cancers. Cell. 2012; 149:979–993. [PubMed: 22608084]
61. McFadden DG, Papagiannakopoulos T, Taylor-Weiner A, et al. Genetic and clonal dissection of murine small cell lung carcinoma progression by genome sequencing. Cell. 2014; 156:1298–311. [PubMed: 24630729]
62. Nixon KC. The Parsimony Ratchet, a New Method for Rapid Parsimony Analysis. Cladistics. 1999; 15:407–414.
63. Lawrence MS, et al. Mutational heterogeneity in cancer and the search for new cancer-associated genes. Nature. 2013; 499:214–8. [PubMed: 23770567]
Figure 1. Barrett’s esophagus has a mutation frequency comparable to many invasive cancers but shows few genomic amplifications.

a) Mutation density (mutations per Mb of sequenced exome) of Barrett’s (BE_{NoDys}), Barrett’s with dysplasia (BE_{Dys}), and esophageal adenocarcinoma (EAC) compared to several other invasive cancers. Mutation density displayed on a logarithmic scale. Box plot shows the median, limited by the 25th (Q1) 75th (Q3) percentiles, with the upper and lower whiskers denoting the extreme most data point within Q_{3} + 1.5*IQR or Q_{1} – 1.5*IQR respectively. N = 133 for prostate, 889 for breast, 14 for BE NoDys, 233 for colorectal, 141 for EAC, 11 for BE Dys, and 401 for lung.

b) “Lego” plots of mutation frequencies across 25 Barrett’s samples (left) and esophageal adenocarcinoma (right). Base substitutions are divided into six categories to represent the six possible base changes (each category represented by a different color). Substitutions are further divided by the 16 possible flanking nucleotides surrounding the mutated base as listed in the corresponding box legend. The inset pie chart indicates the distribution of all mutations for a given base.

c) Mean number of deletions per sample.

d) Mean number of amplifications per sample. * denotes statistically significant difference ($P < 0.05$).
**Figure 2. Paired analysis reveals early-shared TP53 alterations**

a) Tumor suppressor plot showing both mutations (heterozygous or homozygous) and homozygous deletions of 4 commonly altered tumor suppressor genes in the BE samples and their paired EAC. Patients are separated into cases where the BE and EAC samples are clonally unrelated (left) or clonally related (right) with ordering by increasing percent of shared mutations. Orange box indicates alterations that were shared between the two samples whereas the triangles represent alterations private to either the BE or EAC. Sample EAC 71 contained both a shared ARID1A alteration and an exclusive ARID1A alteration. Black bordered boxes denote samples that have undergone genome doubling and * denotes samples suggestive of high-grade dysplasia.

b) Example of evolutionary “tree” where despite only sharing a small percentage of overall mutations a TP53 mutation was found to be one of the early shared events. The lengths of the lines represent that number of mutations in common to this branch according to scale. Thin lines denote alterations with CCF<1.

c) Example of evolutionary “tree” where a CDKN2A mutation occurred late in the Barrett’s sample after the clone that went on to develop into an invasive cancer already split off. * denotes samples suggestive of high-grade dysplasia.
Figure 3. Paired analysis reveals a lack of oncogene amplification in BE samples
Amplification plot showing amplified oncogenes, mutations, and pathways in BE compared
to EAC with the genomic doubling status of samples and the presence or absence of
dysplasia in the BE marked. * denotes samples suggestive of high-grade dysplasia.
Figure 4. Spatial and phylogenic relationship of multiple sampled areas where all samples within a patient share a common set of genomic alterations

a, c, e) Diagram showing relative location of samples isolated for genomic analysis. Samples immediately adjacent to each other were isolated on the same block. Specimens marked with an asterisk did not contain enough information to be properly located. Size of BE and EAC roughly proportional to the reported BE length and tumor size.

b, d, f) Phylogenic trees displaying the relationship of subclones detected in each tissue sample. Branch lengths are proportional to the number of somatic point-mutations occurring on that branch. For mutations detected in a single sample, the thickness of the branch is proportional to the CCF of the mutations in that sample. Starting at the germline, light gray branches represent acquired alterations shared by all samples from a given patient.

b) P1 shows a TP53 mutation shared with all samples as well as a CDK6 amplification only seen in the EAC samples.

d) P4 shows a shared CDKN2A deletion in all samples with TP53 mutation, whole genome doubling (WGD), and oncogene amplification in the highly related high-grade dyplasia and cancer samples.

f) P7 shows a TP53 mutation shared across all samples and KRAS amplification only found in the cancer and a single focus of adjacent HGD. Shaded areas contain the subclones present in the corresponding tissue samples.
Figure 5. Spatial and phylogenetic relationship of multiple sampled areas in patients showing distinct clonal evolution

a, c) Diagram of sample location and diagnosis within the patient’s field of BE. Size of BE and EAC roughly proportional to the reported BE length and tumor size. Specimens marked with an asterisk did not contain enough information to be properly located. b, d) Phylogenetic trees displaying the relationship of subclones detected in each tissue sample. Branch lengths are proportional to the number of somatic point-mutations occurring on that branch. For mutations detected in a single sample, the thickness of the branch is proportional to the CCF of the mutations in that sample. Black circles mark the starting point (germline) as there were no somatic alterations common to all samples. b) P3 shows three distinct clonally unrelated branches. The branch with BE2/LGD1 shows a CDKN2A deletion in the BE and LGD samples only. The green shaded region represents tissue sample EAC1, which contained a mixture of distinct cell populations, subclones 1 and 2. d) P6 shows a region of

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BE tissue unrelated to the other high-grade and cancer samples, all of which share ATM and SMAD2 mutations and WGD.
Figure 6. Tumor suppressor gene alterations are more common in non-genome doubled EAC

(a) Representation of alterations of common tumor suppressors in a larger cohort of EAC samples showing truncating mutations, missense mutations of hotspot site (as determined by presence in the COSMIC repository at least 3 times) and homozygous deletions. Samples are divided into cases that have undergone genome doubling (left) and those that have not (right). The type of mutation identified is represented by the color of the mutation box. 

(b) Expanded analysis of the fraction of genome doubled and non-genome doubled cases with alterations in the given tumor suppressor pathways (genes in pathway with multiple identified alterations listed on right). Statistically significant differences are highlighted. Genes in the individual pathways are also shown in Supplementary Table 5.
Figure 7. Genome doubled EAC contains more frequent amplifications in cell cycle regulators and transcription factors

Amplification plot showing amplified oncogenes, mutations, and pathways in esophageal adenocarcinoma. Samples are divided into cases that have undergone genome doubling (left) and those that have not (right).
Figure 8. Genome doubled EAC shows a distinct pathway of development
Schematic representation showing two general pathways by which BE can develop into EAC. The top model involves the gradual accumulation of tumor suppressor genes followed by the subsequent activation of oncogenes and development of genomic instability. In the bottom model, TP53 inactivation is acquired as an early event. The sample then undergoes genome doubling, leading to genomic instability, aneuploidy, and oncogene amplification.