

1 Pan-cancer whole genome analyses of metastatic solid tumors

2 Peter Priestley^{1,2,*,#}, Jonathan Baber^{1,2,*}, Martijn P. Lolkema^{3,4}, Neeltje Steeghs^{3,5}, Ewart de Bruijn¹,
3 Korneel Duyvesteyn¹, Susan Haidari^{1,3}, Arne van Hoeck⁶, Wendy Onstenk^{1,3,4}, Paul Roepman¹,
4 Charles Shale², Mircea Voda¹, Haiko J. Bloemendal⁷, Vivianne C.G. Tjan-Heijnen⁸, Carla M.L. van
5 Herpen⁹, Mariette Labots¹⁰, Petronella O. Witteveen¹¹, Egbert F. Smit^{3,5}, Stefan Sleijfer^{3,4}, Emile E.
6 Voest^{3,5}, Edwin Cuppen^{1,3,6,#}

7

8 ¹ Hartwig Medical Foundation, Science Park 408, Amsterdam, The Netherlands

9 ² Hartwig Medical Foundation Australia, Sydney, Australia

10 ³ Center for Personalized Cancer Treatment, The Netherlands

11 ⁴ Erasmus MC Cancer Institute, Doctor Molewaterplein 40, Rotterdam, The Netherlands

12 ⁵ Netherlands Cancer Institute/Antoni van Leeuwenhoekhuis, Plesmanlaan 121, Amsterdam, The Netherlands

13 ⁶ Center for Molecular Medicine and Oncode Institute, University Medical Center Utrecht, Heidelberglaan 100,
14 Utrecht, The Netherlands

15 ⁷ Meander Medisch Centrum, Maatweg 3, Amersfoort, The Netherlands

16 ⁸ Maastricht University Medical Center, P. Debyelaan 25, Maastricht, The Netherlands

17 ⁹ Radboud University Medical Center, Geert Grooteplein Zuid 10, Nijmegen, The Netherlands

18 ¹⁰ VU Medical Center, De Boelelaan 1117, Amsterdam, The Netherlands

19 ¹¹ Cancer Center, University Medical Center Utrecht, Heidelberglaan 100, Utrecht, The Netherlands

20

21 * shared first author

22 # corresponding authors: p.priestley@hartwigmedicalfoundation.nl, e.cuppen@hartwigmedicalfoundation.nl

23 Abstract

24 **Metastatic cancer is one of the major causes of death and associated with poor**
25 **treatment efficiency. A better understanding of the characteristics of late stage cancer is**
26 **required to help tailor personalised treatment, reduce overtreatment and improve outcomes.**
27 **Here we describe the largest pan-cancer study of metastatic solid tumor genomes, including**
28 **2,520 whole genome-sequenced tumor-normal pairs, analyzed at a median depth of 106x and**
29 **38x, respectively, and surveying over 70 million somatic variants. Metastatic lesions were**
30 **found to be very diverse, with mutation characteristics reflecting those of the primary tumor**
31 **types, although with high rates of whole genome duplication events (56%). Metastatic lesions**
32 **are relatively homogeneous with the vast majority (96%) of driver mutations being clonal and**
33 **up to 80% of tumor suppressor genes bi-allelically inactivated through different mutational**
34 **mechanisms. For 62% of all patients, genetic variants that may be associated with outcome of**
35 **approved or experimental therapies were detected. These actionable events were distributed**
36 **over the various mutation types (single and multiple nucleotide variants, insertions and**
37 **deletions, copy number alterations and structural variants) underlining the importance of**
38 **comprehensive genomic tumor profiling for cancer precision medicine for advanced cancer**
39 **treatment.**

40 Introduction

41 Metastatic cancer is one of the leading causes of death globally and is a major burden for
42 society despite the availability of an increasing number of (targeted) drugs. Health care costs
43 associated with treatment of metastatic disease are increasing rapidly due to the high cost of novel
44 targeted treatments and immunotherapy, while many patients do not benefit from these approaches
45 with inevitable adverse effects for most patients. Metastatic cancer therefore poses a major challenge
46 for society to balance between individual and societal treatment responsibilities. As cancer genomes

47 evolve over time, both in the highly heterogeneous primary tumor mass and as disseminated
48 metastatic cells^{1,2}, a better understanding of metastatic cancer genomes is crucial to further improve
49 on tailoring treatment for late stage cancers.

50 In recent years, several large-scale whole genome sequencing (WGS) analysis efforts such
51 as TCGA and ICGC have yielded valuable insights in the diversity of the molecular processes driving
52 different types of adult^{3,4} and pediatric^{5,6} cancer and have fueled the promises of genome-driven
53 oncology care⁷. However, these analyses were primarily done on primary tumor material whereas
54 metastatic cancers, which cause the bulk of the disease burden and 90% of all cancer deaths, have
55 been less comprehensively studied at the whole genome level, with previous efforts focusing on
56 tumor-specific cohorts⁸⁻¹⁰ or at a targeted gene panel¹¹ or exome level¹².

57 Here we describe the first large-scale pan-cancer whole-genome landscape of metastatic
58 cancers based on the Hartwig Medical Foundation (HMF) cohort of 2,520 paired tumor and normal
59 genomes from 2,405 patients. The samples have been collected prospectively as fresh frozen
60 biopsies taken from a broad range of metastases (Extended Data Fig. 1b) and blood controls from
61 patients with advanced cancer in a clinical study setup coordinated by the Center for Personalized
62 Cancer Treatment (CPCT) in 41 hospitals in the Netherlands (Supplementary Table 1). All samples
63 were paired with standardized clinical information (Extended Data Fig. 1a). The sample distribution
64 over age and primary tumor types broadly reflects solid cancer incidence in the Western world,
65 including rare cancers (Figure 1a-b).

66 The cohort has been analyzed with uniform and high depth paired-end (2 x 150 bp) whole
67 genome sequencing with a median depth of 106x for tumor samples and 38x for the blood control
68 (Extended Data Fig. 1c). Sequencing data were analyzed for all types of somatic variants using an
69 optimized bioinformatic pipeline based on open source tools (Methods). We identified a total of
70 59,472,629 single nucleotide variants (SNVs), 839,126 multiple nucleotide variants (MNVs),
71 9,598,205 insertions and deletions (INDELs) and 653,452 structural variants (SVs) (Supplementary
72 Table 2). We found that the relative high sequencing depth is important for variant calling sensitivity
73 as downsampling of the tumor sample coverage to ~53x resulted in an average decrease in sensitivity
74 of 10% for SNV, 2% for INDEL, 15% for MNV, and 19% for SV (Extended Data Fig. 2).

75 Here we present a detailed characterization of this unique and comprehensive resource for a
76 better genomic understanding of metastatic cancer.

77 **Mutational landscape of metastatic cancer**

78 We analysed the tumor mutational burden (TMB) of each class of variants per cancer type
79 based on tissue of origin (Fig. 1c-h, Supplementary Table 2). In line with previous studies on primary
80 cancers¹³, we found extensive variation in mutational load of up to 3 orders of magnitude both within
81 and across cancer types.

82 The median SNV counts per sample were highest in skin, predominantly consisting of
83 melanoma (44k) and lung (36k) tumors with ten-fold higher SNV counts than sarcomas (4.1k),
84 neuroendocrine tumors (NET) (3.5k) and mesotheliomas (3.4k). The variation for MNVs was even
85 greater with lung (median=815) and skin (median=764) tumors having five times the median MNV
86 counts of any other tumor type. This can be explained by the well-known mutational impact of UV
87 radiation (CC->TT MNV) and smoking (CC->AA MNV) mutational signatures, respectively (Fig. 1f).
88 Although only di-nucleotide substitutions are typically reported as MNVs, 10.7% of the MNVs involve
89 three nucleotides and 0.6% had four or more nucleotides affected.

90 INDEL counts were typically ten-fold lower than SNVs, with a lower relative rate for skin and
91 lung cancers (Fig. 1d, Extended Data Fig. 3). Genome-wide analysis of INDELs at microsatellite loci
92 identified 60 samples with microsatellite instability (MSI) (Supplementary Table 2), representing 2.4%
93 of all tumors. The highest rates of MSI were observed in central nervous system (CNS) (9.4%), uterus
94 (9.0%) and prostate (6.1%) tumors. For metastatic colorectal cancer lesions we found an MSI
95 frequency of only 4.0%, which is lower than reported for primary colorectal cancer, and in line with
96 better prognosis for patients with localized MSI colorectal cancer, which less often metastasizes¹⁴.

97 Remarkably, 67% of all INDELs in the entire cohort were found in the 60 MSI samples, and 85% of all
98 INDELs in the cohort were found in microsatellites or short tandem repeats. Only 0.33% of INDELs
99 (32k, ~1% of non-microsatellite INDELs) were found in coding sequences, of which the majority (88%)
100 had a predicted high impact by affecting the open reading frame of the gene.

101 The median rate of SVs across the cohort was 193 per tumor, with the highest median counts
102 observed in ovary (415) and esophageal (379) tumors, and the lowest in kidney tumors (71) and NET
103 (56) (Fig. 1h, Supplementary Table 2). Simple deletions were the most commonly observed SV
104 subtype (33% of all SVs) and were the most prevalent in every cancer type except stomach and
105 esophageal tumors which were highly enriched in translocations.

106 Copy number alteration landscape of metastatic cancer

107 Copy number alterations (CNAs) are important hallmarks of tumorigenesis¹⁵. Pan-cancer, the
108 most highly amplified regions in our metastatic cancer cohort contain the established oncogenes such
109 as EGFR, CCNE1, CCND1 and MDM2 (Fig. 2a). Chromosomal arms 1q, 5p, 8q and 20q are also
110 highly enriched in moderate amplification across the cohort each affecting >20% of all samples. For
111 the amplifications of 5p and 8q this is likely related to the common amplification targets of TERT and
112 MYC, respectively. However, the targets of the amplifications on 1q, predominantly found in breast
113 cancers (>50% of samples) and amplifications on 20q, predominantly found in colorectal cancers
114 (>65% of samples), are less clear.

115 We identified some intriguing patterns of recurrent loss of heterozygosity (LOH) caused by
116 CNAs. Overall an average of 23% of the autosomal DNA per tumor has LOH. Unsurprisingly, TP53
117 has the highest LOH recurrence at 67% of samples. Many of the other LOH peaks are also explained
118 by well-known tumor suppressor genes (TSG). However, several clear LOH peaks are observed
119 which cannot easily be explained by known TSG selection, such as one on 8p (57% of samples),
120 which could potentially be the result of the mechanism by which the amplification of 8q, the most
121 commonly amplified part of the genome, is established.

122 There are remarkable differences in LOH between cancer types (Figure 2, Extended Data
123 Fig. 4). For instance, we observed LOH events on the 3p arm in 90% of kidney samples¹⁶ and LOH of
124 the complete chromosome 10 in 72% of CNS tumors (predominantly glioblastoma multiforme¹⁷). Even
125 in the case of the TP53 region on chromosome 17, different tumor types display clearly different
126 patterns of LOH. Ovarian cancers exhibit LOH of the full chromosome 17 in 75% of samples whereas
127 in prostate cancer, which also has 70% LOH for TP53, this is nearly always caused by highly focal
128 deletions.

129 Unlike LOH events, homozygous deletions are nearly always restricted to small chromosomal
130 regions. Not a single example was found in which a complete autosomal arm was homozygously
131 deleted. Homozygous deletions of genes are surprisingly rare as well: we found only 4,915 autosomal
132 events (average of two per tumor) where one or multiple consecutive genes are fully or partially
133 homozygously deleted. In 46% of these events a putative TSG was deleted. The scarcity of
134 passenger homozygous deletions underlines the fact that despite widespread copy number
135 alterations in metastatic tumors, the vast majority of genes or gross chromosomal organization likely
136 remain essential for tumor cell survival. Chromosome Y is a special case and is deleted in 35% of all
137 male tumor genomes, although this varies significantly between cancer types from 5% in CNS tumors
138 to as high as 68% in biliary tumors.

139 An extreme form of copy number changes can be caused by whole genome duplication
140 (WGD). We found WGD events in 56% of all samples ranging between 17% in CNS to 80% in
141 esophageal tumors (Fig. 2d,e). This is much higher than reported previously for primary tumors (25%-
142 37%)^{18,19} and also higher than estimated from panel-based sequencing analyses of advanced tumors
143 (30%)²⁰. Ploidy levels, in combination with tumor purity data, are essential for correct interpretation of
144 the measured raw SNV and INDEL frequencies, e.g. to discriminate causal bi-allelic inactivation of
145 TSG from heterozygous passenger events. Hence determining the WGD status of a tumor is highly
146 relevant for diagnostic applications. Furthermore, WGD has previously been found to correlate with a

147 greater incidence of cancer recurrence for ovarian cancer¹⁹ and has been associated with poor
148 prognosis across cancer types, independently of established clinical prognostic factors²⁰.

149 **Discovery of novel significantly mutated genes**

150 To identify significantly mutated genes (SMGs) potentially specific for metastatic cancer, we
151 used the dNdScv approach²¹ with strict cutoffs ($q < 0.01$) for the pan-cancer and tumor-type specific
152 datasets. In addition to reproducing previous results on cancer drivers, a few novel genes were
153 identified (**Extended Data Fig. 5, Supplementary Table 3**). In the pan-cancer analyses we found only a
154 single novel SMG, which was not either present in the curated COSMIC Cancer Gene Census or
155 found by Martincorena et al²¹. This gene, MLK4 (q -value = $2e-4$), is a mixed lineage kinase that
156 regulates the JNK, P38 and ERK signaling pathways and has been reported to inhibit tumorigenesis in
157 colorectal cancer²². In addition, in our tumor type-specific analyses, we identified a novel breast
158 cancer-specific SMG - ZFPM1 (also known as Friend of GATA1 (FOG1), q -value = $8e-5$), a zinc-finger
159 transcription factor protein without clear links with cancer. Nonetheless, we found six unique
160 frameshift variants (all inactivated biallelically) and three nonsense variants, which suggests a driver
161 role for this gene in breast cancer.

162 Our cohort also lends support to some prior SMG findings. In particular, eight significantly
163 mutated putative TSG in the HMF cohort were also found by Martincorena et al²¹ - GPS2 (pan-cancer,
164 $q = 1e-5$ & breast, $q = 2e-3$), SOX9 (colorectal & pan-cancer, $q = 0$), TGIF1 (pan-cancer, $q = 3e-3$ &
165 colorectal $q = 6e-3$), ZFP36L1 (urinary tract $q = 3e-4$, pan-cancer $q = 9e-4$) and ZFP36L2 (colorectal &
166 pan-cancer, $q = 0$), HLA-B (lymphoid, $q = 5e-5$), MGA (pan-cancer, $q = 4e-03$), KMT2B (skin, $q = 3e-3$) and
167 RARG (urinary tract $8e-4$). None of these genes are currently included in the COSMIC Cancer Gene
168 Census²³. ZFP36L1 and ZFP36L2 are of particular interest as these genes are zinc-finger proteins
169 that normally play a repressive regulatory role in cell proliferation, presumably through a cyclin D
170 dependent and p53 independent pathway²⁴. ZFP36L2 is also independently found as a significantly
171 deleted gene in our cohort, most prominently in colorectal and prostate cancers.

172 We also searched for genes that were significantly amplified or deleted (**Supplementary Table**
173 **4**). CDKN2A and PTEN were the most significantly deleted genes overall, but many of the top genes
174 involved common fragile sites (CFS) particularly FHIT and DMD, deleted in 5% and 4% of samples,
175 respectively. The role of CFS in tumorigenesis is unclear and they are frequently treated as
176 passenger mutations reflecting localized genomic instability²⁵. However, the uneven distribution of the
177 deletions across cancer types may indicate that some of these could be genuine tumor-type specific
178 cancer drivers. For example, we find deletions in DMD to be highly enriched in esophageal tumors
179 (38% deleted), GIST (Gastro-Intestinal Stromal Tumors; 24%) and pancreatic NET (41%), which is
180 consistent with a recent study that indicated DMD as a TSG in cancers with myogenic programs²⁶.
181 We also identified several significantly deleted genes not reported previously, including MLLT4 (13
182 samples) and PARD3 (9 samples).

183 Unlike homozygous deletions, amplification peaks tend to be broad and often encompass
184 large number of genes, making identification of the amplification target challenging. However, SRY-
185 related high-mobility group box 4 gene (SOX4, 6p22.3) stands out as a significantly amplified single
186 gene peak (26 amplifications) and is highly enriched in urinary tract cancers (19% of samples highly
187 amplified) (**Extended Data Fig. 4**). SOX4 is known to be over-expressed in prostate, hepatocellular,
188 lung, bladder and medulloblastoma cancers with poor prognostic features and advanced disease
189 status and is a modulator of the PI3K/Akt signaling²⁷.

190 Also notable was a broad amplification peak of 10 genes around ZMIZ1 at 10q22.3 (32
191 samples) which has not previously been reported. ZMIZ1 is a member of the Protein Inhibitor of
192 Activated STAT (PIAS)-like family of coregulators and is a direct and selective cofactor of Notch1 in T-
193 cell development and leukemia²⁸. CDX2, previously identified as an amplified lineage-survival
194 oncogene in colorectal cancer²⁹, is also highly amplified in our cohort with 20 out of 22 amplified
195 samples found in colorectal cancer, representing 5.4% of all colorectal samples.

196 Driver mutation catalog

197 We created a comprehensive catalog of all cancer driver mutations across all tumors in our
198 cohort and all variant classes similar as described before³⁰ (N. Lopez, personal communication). To
199 do this, we combined our SMG discovery efforts with those from Martincorena et al.²¹ and a panel of
200 well known cancer genes (Cosmic Curated Genes)³¹, and searched for fusions and TERT promoter
201 mutations in our cohort. Accounting for the proportion of SNV and INDELS estimated by dNdScv to be
202 passengers, we found 13,433 driver events among the 20,352 identified mutations in the combined
203 gene panel (**Supplementary table 5**). This includes 7,423 coding mutation drivers, 615 non-coding
204 point mutation drivers, 2,715 homozygous deletions (25% of which are in common fragile sites), 2,393
205 focal amplifications and 287 fusion events. To facilitate analysis of variants of unknown significance
206 (VUS) at a per patient level, we calculated a sample specific likelihood for each point mutation to be a
207 driver taking into account the TMB of the sample as well as the biallelic inactivation status of the gene
208 for TSG and hotspot positions in oncogenes. Of note, only 52% of point mutations in driver genes
209 were predicted to be genuine driver events. Predictions of pathogenic variant overlap with known
210 biology, e.g. clustering of benign missense variants in the 3' half of the APC gene (**Extended Data Fig.**
211 **6b**) fits with the absence of FAP-causing germline variants in this part of the gene³².

212 Overall, the catalog matches previous inventories of cancer drivers. TP53 (52% of samples),
213 CDKN2A (21%), APC (16%), PIK3CA (16%), KRAS (14%), PTEN (13%) and TERT (12%) were the
214 most common driver genes together making up 25% of all the driver mutations in the catalog (**Fig. 3**).
215 However, all of the ten most prevalent driver genes in our cohort were reported at a higher rate than
216 for primary cancers³³, which may reflect the more advanced disease state. AR and ESR1 were found
217 as outliers with driver mutations in 44% of prostate and 18% of breast cancers, respectively. Both
218 genes are linked to resistance to hormonal therapy, a common treatment for these tumor types, and
219 have been previously reported as enriched in advanced metastatic cancer¹¹ but are identified at
220 higher rates in this study.

221 Looking at a per patient level, the mean number of mutated driver genes per patient was 5.6
222 (including any type of mutational event), with highest rate in urinary tract tumors (mean rate = 8.0) and
223 the lowest in NET (mean rate = 2.8) (**Fig. 4**). Esophageal and stomach tumors also had highly
224 elevated driver counts, largely due to a much higher rate of deletions in CFS genes (mean rate = 1.5
225 for stomach, 1.7 for esophageal) compared to other cancer types (pan-cancer mean rate = 0.3).
226 Fragile sites aside, the differential rates of drivers between cancer types in each variant class do
227 correlate with the relative mutational load, with the exception of skin cancers, which have a lower than
228 expected number of SNV drivers (**Extended Data Fig. 2**).

229 In 98.5% of all samples at least one driver mutation was found. Of the 36 samples in which no
230 tumor driver mutation was identified, 18 were NET of the small intestine (representing 46.5% of all
231 patients in this category). This likely indicates that small intestine NETs have a distinct set of drivers
232 that are not captured yet in any of the cancer gene resources used and are also not prevalent enough
233 in our relatively small NET cohort to be detected as significant.

234 The number of amplified driver genes varied significantly between cancer types with highly
235 elevated rates per sample in breast cancer (mean = 2.0), esophageal, urinary tract and stomach (all
236 mean = 1.7) cancers and nearly no amplification drivers in kidney cancer (mean = 0.1) and none in
237 the mesothelioma cohort (**Extended Data Fig. 7a**). In tumor types with high rates of amplifications,
238 these amplifications are generally found across a broad spectrum of oncogenes (**Extended Data Fig.**
239 **7b**), suggesting there are mutagenic processes active in these tissues that favor amplifications, rather
240 than tissue-specific selection of individual driver genes. AR and EGFR are notable exceptions, with
241 highly selective amplifications in prostate, and in CNS and lung cancers, respectively, in line with
242 previous reports^{17,34,35}. Intriguingly, we also found two-fold more amplification drivers in samples with
243 WGD (**Extended Data Fig. 7c**) despite amplifications being defined as relative to the average sample
244 ploidy.

245 Analysis of known fusion drivers and promiscuous fusion partners identified 175 in-frame
246 fusion in genic regions and 92 cis-activating fusions involving repositioning of regulatory elements,
247 and 20 in-frame intragenic deletions where one or more exons were deleted. ERG (89 samples),

248 BRAF(17 samples), ERBB4 (16 samples), ALK(12 samples), NRG1(9 samples) and ETV4 (7
249 samples) were the most commonly observed 3' partners together making up more than half of the
250 fusions. 77 of the 89 ERG fusions were TMPRSS2-ERG affecting 37% of all prostate cancer samples
251 in the cohort. 153 fusion pairs were not previously recorded in CGI, OncoKb, COSMIC or CIVIC^{31,36-38}.
252 A novel recurrent KMT2A-BCOR fusion was observed in 2 samples (sarcoma and stomach cancer)
253 and there were also 5 recurrent localized fusions resulting from adjacent gene pairs: NSD1-ZNF346
254 (3 samples), FGFR2-ATE1 (2 samples), KMT2A-BCOR(2 samples), AGM1-ETV1 (2 samples), and
255 BCR-GNAZ (2 samples).
256 Only promoter mutations in TERT were included in the study due to the current lack of robust
257 evidence for other recurrent oncogenic non-coding mutations³⁹. A total of 257 variants were found at 5
258 known recurrent variant hotspots¹¹, and included in the driver catalog.

259 **Oncogene hotspots and novel activating variants**

260 We found that the 70% of driver mutations in oncogenes occur at or within 5 nucleotides of
261 already known pathogenic mutational hotspots (Extended Data Fig. 6a). In the six most prevalent
262 oncogenes (KRAS, PIK3CA, BRAF, NRAS, CTNNB1 & ESR1) the rate was 96% (Fig. 5).
263 Furthermore, in many of the key oncogenes, we also found several likely activating but non-canonical
264 variants near known mutational hotspots (Fig. 5). For example, we found activating MNVs in the well
265 known BRAF V600 hotspot (22 cases), but also novel non-hotspot MNVs in KRAS (8 cases) and
266 NRAS (4 cases) (Extended Data Fig 6b).

267 In-frame indels were even more striking, since despite being exceptionally rare overall (mean
268 = 1.7 per sample), we found an excess in known oncogenes including PIK3CA (19 cases), ERBB2
269 (10 cases) and BRAF(8 cases) frequently occurring at or near known hotspots. Notably, many of the
270 in-frame indels are enriched in specific tumor types. For instance, all 18 KIT in-frame indels were
271 found in sarcomas, 6 out of 8 MUC6 in-frame indels in esophageal tumors, and 6 of 10 ERBB2 in-
272 frame indels in lung tumors. Finally, we identified 10 in-frame indels in FOXA1, which are highly
273 enriched in prostate cancer (7 of 10 cases) and clustered in two locations that were not previously
274 associated with pathogenic mutations⁴⁰.

275 In CTNNB1 we identified an interesting novel recurrent in-frame deletion of the complete exon
276 3 in 12 samples, 9 of which are colorectal cancers. Surprisingly, these deletions were homozygous
277 but suspected to be activating as CTNNB1 normally acts as an oncogene in the WNT/beta-catenin
278 pathway and none of these nine colorectal samples had any APC driver mutations.

279 **Biallelic tumor suppressor gene inactivation**

280 Our results strongly support the Knudson two-hit hypothesis⁴¹ for tumor suppressor genes
281 with 80% of all TSG drivers explained by biallelic inactivation (i.e. either by homozygous deletion
282 (32%), multiple somatic point mutations (7%), or a point mutation in combination with LOH (41%)).
283 This rate is the highest observed in any large-scale cancer WGS study. For many key tumor
284 suppressor genes the biallelic inactivation rate is almost 100% (more specifically: TP53 (93%),
285 CDKN2A (97%), RB1 (94%), PTEN (92%) and SMAD4 (96%); Fig. 3b), suggesting that biallelic
286 inactivation of these genes is a strict requirement for metastatic cancer.

287 Other prominent TSGs, however, have lower biallelic rates, including ARID1A (55%), KMT2C
288 (49%) and ATM (49%). It is unclear whether we systematically missed the second hit in these cases,
289 as this could potentially be mediated through non-mutational epigenetic mechanisms⁴² or unknown
290 pathogenic germline variants, or if these genes impact on tumorigenesis by a haploinsufficiency
291 mechanism⁴³.

292 **Clonal and subclonal variants**

293 To obtain insight into ongoing tumor evolution dynamics, we examined the clonality of all
294 variants. Surprisingly, only 6.5% of all SNV, MNV & INDELS across our cohort and just 3.7% of our

295 driver point mutations were found to be subclonal (Fig. 6). The low proportion of samples with
296 subclonal variants could be partially due to the detection limits of the sequencing approach,
297 particularly for low purity samples, even with our relatively high WGS sequencing depth. However,
298 even for samples with purities higher than 80% the proportion of subclonal variants only reaches
299 10.2% (Fig. 6c). This relatively high degree of tumor homogeneity may be in part attributed to the fact
300 that nearly all biopsies were obtained by a core needle biopsy, which results in highly localized
301 sampling, but is nevertheless much lower compared to previous observations in primary cancers².

302 In the 111 patients with independently collected repeat biopsies from the same patient
303 (Supplementary Table 6) we found 11% of all SNVs to be subclonal. Whilst 76% of clonal variants
304 were shared between biopsies, less than 30% of the subclonal variants were shared. Together, the
305 low rate of subclonal variants, and the observation that a very high proportion are private to a local
306 metastasis, suggest a model where individual metastatic lesions are dominated by a single clone at
307 any one point in time and that more limited tumor evolution and subclonal selection takes place after
308 distant metastatic seeding.

309 Co-occurrence of Drivers

310 We examined the pairwise co-occurrence of driver gene mutations per cancer type and found
311 ten combinations of genes that were significantly mutually exclusively mutated, and ten combinations
312 of genes which were significantly co-occurently mutated (excluding pairs of genes on the same
313 chromosome which are frequently co-amplified or co-deleted) (Fig. 7). The 20 significant findings
314 include previously reported co-occurrence of mutated DAX|MEN1 in pancreatic NET ($q=0.0007$), and
315 CDH1|SPOP in prostate tumors ($q=0.0005$), as well as negative associations of mutated genes within
316 the same signal transduction pathway such as KRAS|BRAF ($q=4e-4$) & KRAS|NRAS ($q=0.009$) in
317 colorectal cancer, BRAF|NRAS in skin cancer ($q=6e-12$), CDKN2A|RB1 in lung cancer ($q=8e-5$) and
318 APC|CTNNB1 in colorectal cancer ($q=8e-6$). APC is also strongly negatively correlated with BRAF
319 ($q=1e-4$) and RNF43 ($q=2e-5$) which together are characteristic of the serrated molecular subtype of
320 colorectal cancers⁴⁴. We also found that SMAD2|SMAD3 are highly positively correlated in colorectal
321 cancer ($q=0.02$), mirroring a result reported previously in a large cohort of colorectal cancers⁴⁵.

322 In breast cancer, we found a number of significant novel relationships, including a positive
323 relationship for GATA3|VMP1 ($q=1e-4$) and FOXA1|PIK3CA ($q=2e-3$), and negative relationships for
324 ESR1|TP53 ($q=9e-4$) and GATA3|TP53 ($q=2e-3$).

325 Actionability

326 We analyzed opportunities for biomarker-based treatment for all patients by mapping driver
327 events to three clinical annotation databases: CGI³⁸, CIVIC³⁶ and OncoKB³⁷. In 1,485 patients (62%)
328 at least one 'actionable' event was identified (Supplementary Table 7). While these numbers are in
329 line with results from primary tumors³⁰, longitudinal studies will be required to conclude if genomic
330 analyses for therapeutic guidance should be repeated when a patient experiences progressive
331 disease. Half of the patients with an actionable event (31% of total) contained a biomarker with a
332 predicted sensitivity to a drug at level A (approved anti-cancer drugs) and lacked any known
333 resistance biomarkers for the same drug (Fig. 8). In 13% of patients the suggested therapy was a
334 registered indication, while in 18% of cases it was outside the labeled indication. In a further 31% of
335 patients a level B (experimental therapy) biomarker was identified. The actionable biomarkers
336 spanned all variant classes including 1,815 SNVs, 48 MNVs, 195 indels, 745 CNAs, 68 fusion genes
337 and 60 patients with microsatellite instability.

338 Tumor mutation burden is an important emerging biomarker for response to immune
339 checkpoint inhibitor therapy⁴⁶ as it is a proxy for the amount of neo-antigens in the tumor cells. For
340 NSCLC it has been shown in at least 2 subgroup analyses of large phase III trials that both PFS and
341 OS are significantly improved with first line immunotherapy as compared to chemotherapy for patients
342 whose tumors have a TMB >10 mutations per Mb^{47,48}. Although various clinical studies based on this
343 parameter are currently emerging, TMB was not yet included in the above actionability analysis.

344 However, when applying the same cut-off to all samples in our cohort, an overall 18% of patients
345 would qualify, varying from 0% for liver, mesothelioma and ovarian cancer patients to more than 50%
346 of lung and skin cancer patients (Extended Data Fig. 3b).

347 Discussion

348 Genomic testing of tumors faces numerous challenges in meeting clinical needs, including i)
349 the interpretation of variants of unknown significance (VUS), ii) the steadily expanding universe of
350 actionable genes, often with an increasingly small fraction of patients affected (e.g. NRG-1⁴⁹ and
351 NTRK fusions⁵⁰ in less than 2% of all patients), and iii) the development of advanced genome-derived
352 biomarkers such as tumor mutational load, DNA repair status and mutational signatures. Our results
353 demonstrate in several ways that WGS analyses of metastatic cancer can provide novel and relevant
354 insights and be instrumental in addressing some of these key challenges in cancer precision
355 medicine.

356 First, our systematic and large-scale pan-cancer analyses on metastatic cancer tissue
357 allowed for the identification of several novel (cancer type-specific) cancer drivers and mutation
358 hotspots. Second, the driver catalog analyses can be used to alleviate the problem of VUS
359 interpretation³⁰ by leveraging previously identified pathogenic mutations (accounting for more than
360 2/3rds of oncogenic point-mutation drivers) and by careful analysis of the biallelic inactivation of
361 putative TSG which accounts for over 80% of TSG drivers in metastatic cancer. Third, we
362 demonstrate the importance of accounting for all types of variants, including large scale genomic
363 rearrangements (via fusions and copy number alteration events), which account for more than half of
364 all drivers, but also activating MNV and INDELS which we have shown are commonly found in many
365 key oncogenes. Fourth, we have shown that using WGS, even with very strict variant calling criteria,
366 we could find driver variants in more than 98% of all metastatic tumors, including putatively actionable
367 events in a clinical and experimental setting for up to 62% of patients.

368 Although we did not find metastatic tumor genomes to be fundamentally different to primary
369 tumors in terms of the mutational landscape or genes driving advanced tumorigenesis, we described
370 characteristics that could contribute to therapy responsiveness or resistance in individual patients. In
371 particular we showed that WGD is a more pervasive element of tumorigenesis than previously
372 understood affecting over half of all metastatic cancers. We also found metastatic lesions to be less
373 heterogeneous than primary tumors, supporting a model for metastasis of clonal seeding followed by
374 rapid expansion.

375 It should be noted that differences between WGS cohorts should be interpreted with some
376 caution as inevitable differences between experimental and computational approaches may impact on
377 observations and can only be addressed in longitudinal studies including the different stages of
378 disease. Furthermore, the HMF cohort includes a mix of treatment-naive metastatic patients and
379 patients who have undergone (extensive) prior systemic treatments. While this may impact on specific
380 tumor characteristics, it also provides opportunities for studying treatment response and resistance as
381 this data is recorded in the studies.

382 Finally, we believe the resource described here is a valuable complementary resource to
383 comparable whole genome sequencing-based resources of primary tumors in advancing fundamental
384 and translational cancer research. Therefore, all non-privacy sensitive data is publicly available
385 through a local interface developed by ICGC⁵¹ (work in progress) and all other data is made freely
386 available for scientific research by a controlled access mechanism (see
387 www.hartwigmedicalfoundation.nl for details).

388

389 Acknowledgements

390 We thank the Hartwig Foundation and Barcode for Life for financial support of clinical studies and
391 WGS analyses. Development of the data portal was supported by a grant from KWF
392 Kankerbestrijding (HMF2017-8225, GENONCO). We are particularly grateful to all patients, nurses
393 and medical specialists for their essential contributions making this study possible. We would like to
394 specifically thank Hans van Snellenberg for operational management of the Hartwig Medical
395 Foundation. We would like to thank Stefan Willems, Wendy de Leng, Alexander Hoischen and
396 Winand Dinjens for support with pathology assessments and mutation validations and Jeroen de
397 Ridder, Wigard Kloosterman and Harmen van de Werken for critically reading the manuscript.

398

399

400 Figure Legends

401 **Figure 1: Mutational load of metastatic cancer per tumor type**

402 a) The number of samples of each tumor type cohort. Tumor types are ranked from lowest to highest
403 overall mutation burden (TMB)
404 b) Violin plot showing age distribution of each tumor type with 25th, 50th and 75th percentiles marked.
405 c)-d) cumulative distribution function plot of mutational load for each tumor type for SNV and MNV (c)
406 and INDEL and SV (d). The median for each cohort is indicated with a vertical line.
407 e)-h) Mutational context or variant subtype per individual sample for each of (e) Single Nucleotide
408 Variant (SNV), (f) Multi Nucleotide Variant (MNV), (g) INsertion/DELetion (INDEL), (h) Structural
409 Variant (SV). Each column chart is ranked within tumor type by mutational load in that variant class.
410 MNVs are classified by the dinucleotide substitution with NN referring to any dinucleotide
411 combination. SVs are classified by type: INV = inversion, DEL = deletion, DUP = tandem duplication,
412 TRL = translocation, INS = insertion.

413

414 **Figure 2: Copy number landscape of metastatic cancer**

415 a)-c) Proportion of samples with amplification and deletion events by genomic position per cohort -
416 pan-cancer (a), kidney (b) and central nervous system (CNS) (c). Inner ring shows % of tumors with
417 homozygous deletion (orange), LOH and significant loss (copy number < 0.6x sample ploidy - dark
418 blue) and near copy neutral LOH (light blue). Outer ring shows % of tumors with high level
419 amplification (>3x sample ploidy - orange), moderate amplification (>2x sample ploidy - dark green)
420 and low level amplification (>1.4x amplification - light green). Scale on both rings is 0-100% and
421 inverted for the inner ring. The most frequently observed high-level gene amplifications (black text)
422 and homozygous deletions (red text) are shown.
423 d) Proportion of tumors with a whole genome duplication event (dark blue) grouped by tumor type.
424 e) Average sample ploidy distribution over the complete cohort. Samples with a WGD event (true) are
425 shown in darker blue.

426

427 **Figure 3: Most prevalent driver genes in metastatic cancer**

428 Thirty most prevalent mutated oncogenes (a) and tumor suppressor genes (TSG) (b). From left to
429 right, the heatmap shows the % of samples in each cancer type which are found to have each gene
430 mutated; absolute bar chart shows the pan-cancer % of samples with the given gene mutated; relative
431 bar chart shows the breakdown by type of alteration. For TSG only the % of samples with a driver in
432 which the gene is found biallelically inactivated is also shown.

433

434 **Figure 4: Drivers per sample by tumor type**

- 435 a) Violin plot showing the distribution of number of drivers per sample grouped by tumor type. Black
436 dot indicates the mean value.
437 b) Relative bar chart showing the breakdown per cancer type of the type of alteration.

438

439 **Figure 5: Oncogenic Hotspots**

440 Count of driver point mutations by variant type. Known pathogenic mutations curated from external
441 databases are categorized as hotspot mutations. Mutations within 5 bases of a known pathogenic
442 mutation are shown as near hotspot and all other mutations are shown as non-hotspot.

443

444 **Figure 6: Subclonality**

- 445 a) Count of samples per tumor purity bucket
446 b) Violin plot showing the percentage of point mutations which are subclonal in each purity bucket.
447 Black dot indicates the mean.
448 c) Percentage of driver point mutations that are subclonal in each purity bucket.

449

450 **Figure 7: Driver co-occurrence**

- 451 a) Driver pairs which are significantly positively (on the right) or negatively (on the left) correlated in
452 individual tumor types sorted by q-value. Color indicates tumor type as depicted below the chart.

453

454 **Figure 8: Actionability**

455 Percentage of samples in each cancer type with an actionable mutation based on data in CGI, CIViC
456 and OncoKB knowledgebases. Level 'A' represents presence of biomarkers with either an approved
457 therapy or guidelines and level B represents biomarkers having strong biological evidence or clinical
458 trials indicating they are actionable. On label indicates treatment registered by federal authorities for
459 that tumor type, while off-label indicates a registration for other tumor types.

460

461 **Extended Data Figures and Tables**

462

463 **Extended Data Figure 1: Hartwig sample workflow, biopsy locations and sequence coverage**

- 464 A) Sample workflow from patient to high-quality WGS data. A total of 4,018 patients were enrolled in
465 the study between April 2016 and April 2018. For 9% of patients no blood and/or biopsy material was
466 obtained, mostly because conditions of patients prohibited further study participation. Up to 4 fresh-
467 frozen biopsies per patient were received, which were sequentially analyzed to identify a biopsy with
468 more than 30% tumor cellularity as determined by routine histology assessment. For 859 patients no
469 suitable biopsy was obtained and 2,796 patients were further processed for WGS. 44 and 29 samples
470 failed in either DNA isolation or library preparation and raw WGS data quality QC, respectively. For an
471 additional 385 samples the WGS data was of good quality, but the tumor purity determination based
472 on WGS data (PURPLE) was less than 20% making reliable and comprehensive somatic variant
473 calling and were therefore excluded. Eventually, 2,338 tumor-normal sample pairs with high-quality
474 WGS data were obtained, which were supplemented with 182 pairs from pre-April 2016, adding up to
475 2,520 tumor normal pairs that were included in this study.
476 B) Breakdown of cohort by biopsy location. Tumor biopsies were taken from a broad range of
477 locations. Primary tumor type is shown on the left and the biopsy location on the right.
478 c) Distribution of sample sequencing depth for tumor and blood reference.

479

480 **Extended Data Figure 2: Impact of downsampling on variant calling**

481 Comparison of variant calling of 10 randomly selected samples at normal depth and 50%
482 downsampled for purity (a), SNV counts (b), SV counts (c), Ploidy (d), MNV counts (e) and INDEL

483 counts (f). For the panels B, C, E and F, the black dots represent the % reduction per sample of
484 counts (right axis) and the dotted line represents the average % reduction across all tested samples.

485

486 **Extended Data Figure 3: Mutational load, genome wide analyses and drivers**

487 a) Proportion of samples by cancer type classified as microsatellite instable (MSI)

488 b) Proportion of samples with a high mutational burden (TMB > 10 SNV / Mb)

489 c) Scatter plot of INDEL vs SNV mutational load. MSI and 'high TMB' thresholds are indicated.

490 d)-f) Mutational load vs driver rate for SNV (d), INDEL (e) and SV (f) grouped by cancer type. MSI
491 samples were excluded.

492

493 **Extended Data Figure 4: Copy Number profile per cancer types**

494 Proportion of samples with amplification and deletion events by genomic position per cancer type.

495 Inner ring shows % of tumors with homozygous deletion (red), LOH and significant loss (copy number
496 < 0.6x sample ploidy - dark blue) and near copy neutral LOH (light blue). Outer ring shows % of
497 tumors with high level amplification (>3x sample ploidy - orange), moderate amplification (>2x sample
498 ploidy - dark green) and low level amplification (>1.4x amplification - light green). Scale on both rings
499 are 0-100%, inverted for inner ring. The most frequently observed high level gene amplifications
500 (black text) and homozygous deletions (red text) are labelled.

501

502 **Extended Data Figure 5: Significantly mutated genes**

503 Tile chart showing genes found to be significantly mutated per cancer type cohort and pan-cancer
504 using dNdScv. Gene names marked in orange are also significant in Martincorena et al²¹, but not
505 found in COSMIC curated or census. Gene names marked in red are novel in this study.

506

507 **Extended Data Figure 6: Coding mutation profiles by driver gene**

508 Location and driver classification of all coding mutations (SNVs and indels) in oncogenes (a) and
509 tumor suppressor genes (TSG) (b) in the driver catalog. The lollipops on the chart show the location
510 (coding sequence coordinates) and count of mutations for all candidate drivers. Dotted lines show the
511 count of all variants at each location while solid lines represent the sum of the driver likelihoods
512 assigned to that specific variant. TSG variants are shaded by their biallelic inactivation status. 'Mixed'
513 means that at least 2 samples with the variant have different biallelic statuses. The bars on the right of
514 the chart show the estimated number of drivers (calculated as the sum of driver likelihoods) and
515 passenger variants in the gene by cancer type.

516

517 **Extended Data Figure 7: Amplifications**

518 a) Mean rate of amplification drivers per cancer type

519 b) Number of amplification drivers per gene showing the breakdown by cancer type

520 c) Mean rate of drivers per variant type for samples with and without WGD.

521

522 **Supplementary Table 1:** Overview of contributing organizations and local principal investigators.

523

524 **Supplementary Table 2:** Overview of cohort and sample characteristics

525

526 **Supplementary Table 3:** Pan-cancer and cancer type-specific dNdScv results

527

528 **Supplementary Table 4:** Recurring amplifications (a) and deletions (b) and associated target genes

529

530 **Supplementary Table 5:** Driver catalog

531

532 **Supplementary Table 6:** Overview of patients with multiple biopsies

533

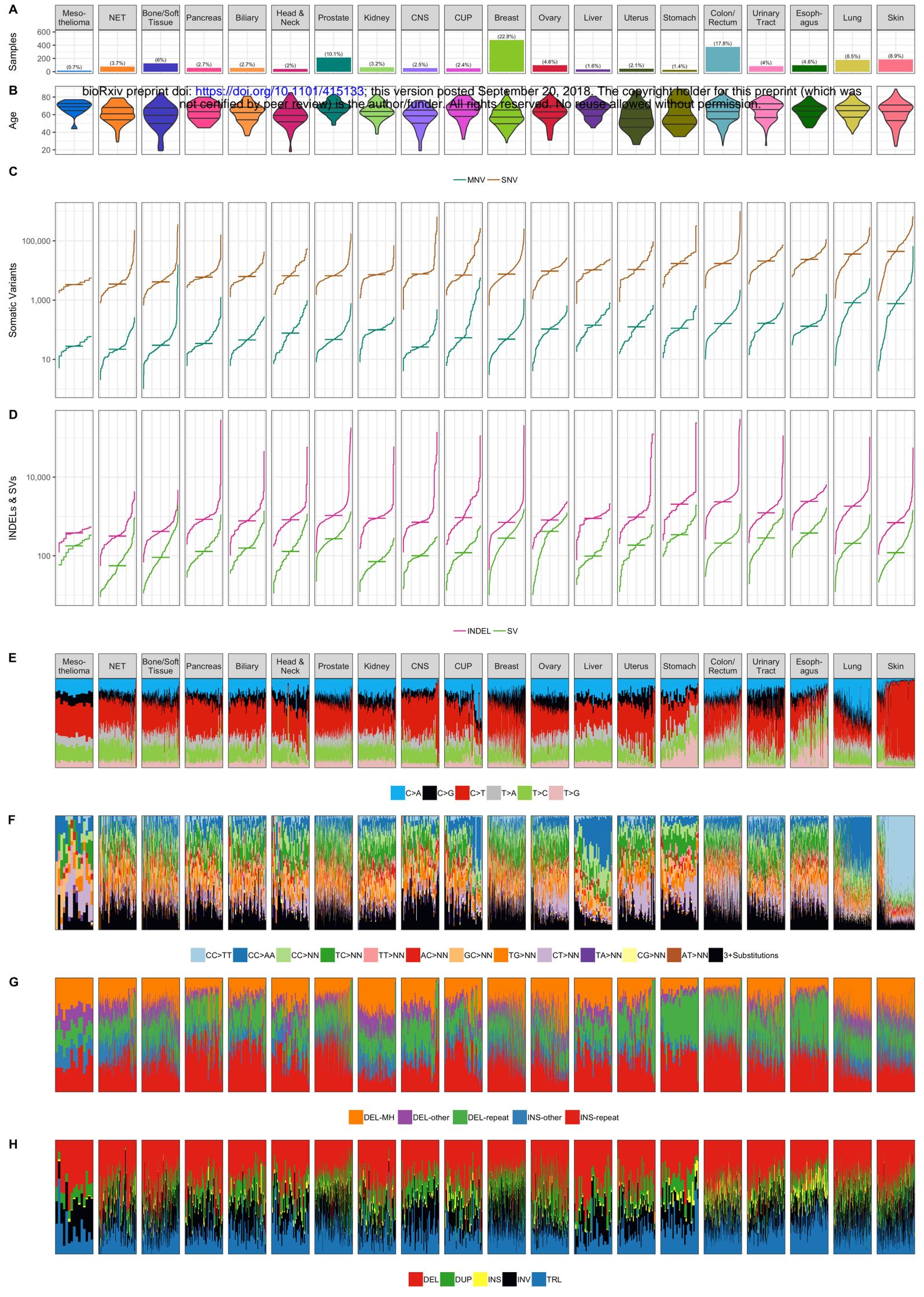
534 **Supplementary Table 7:** Actionable mutations

535 References

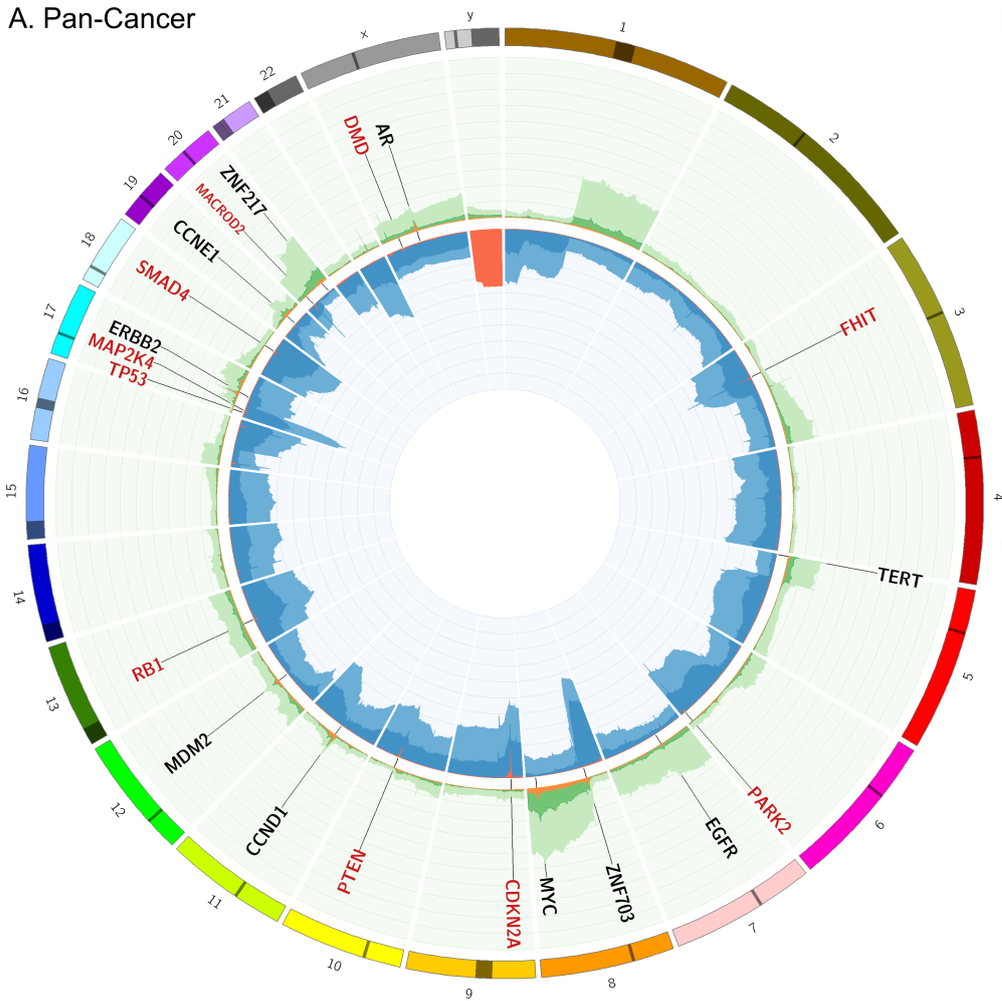
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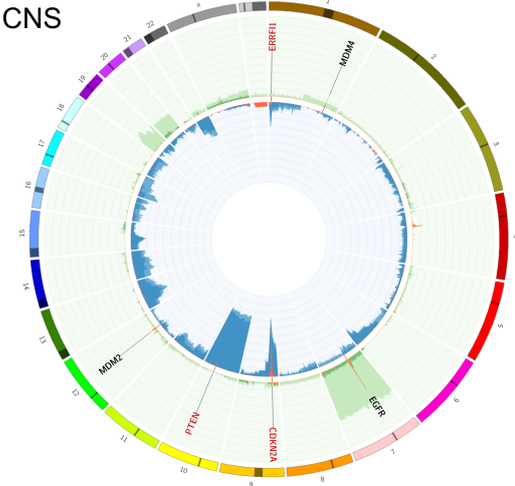
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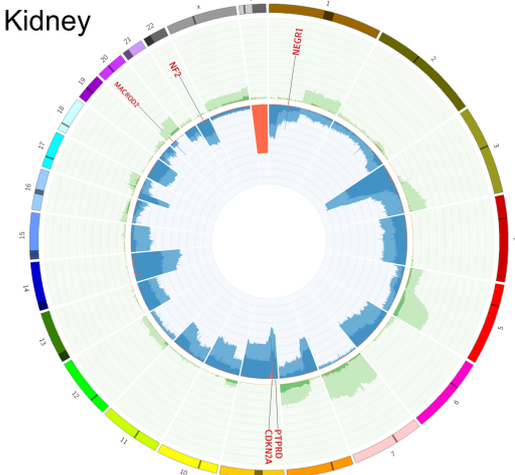
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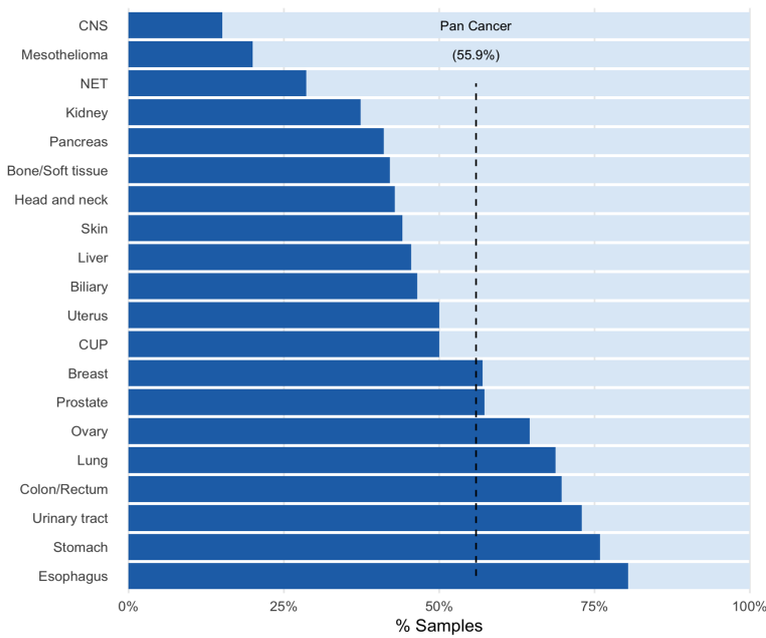
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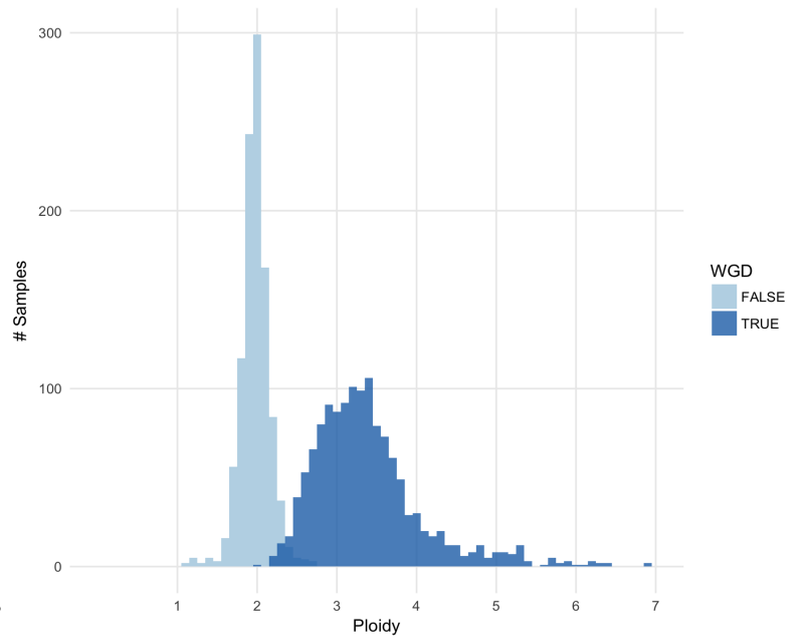
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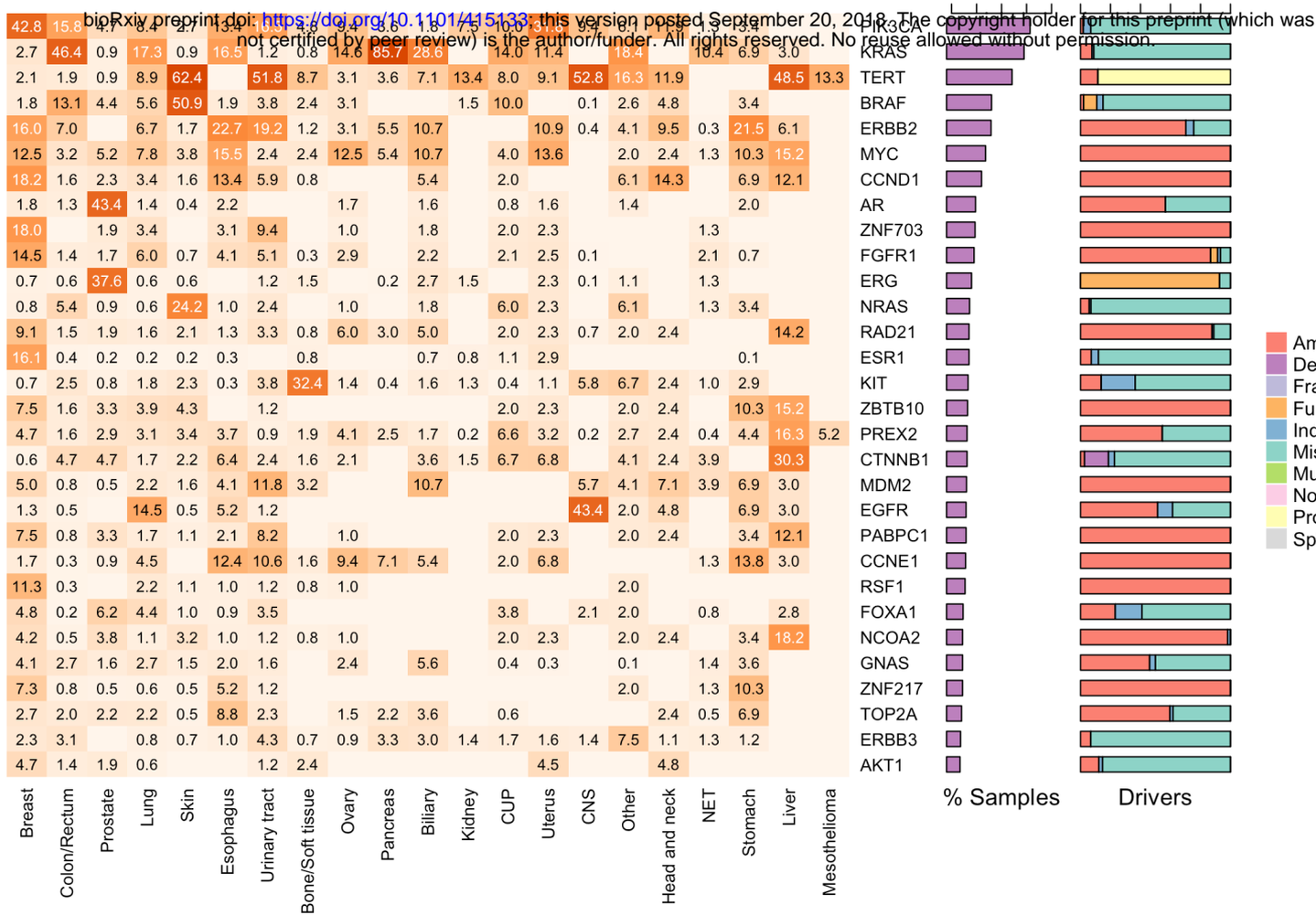
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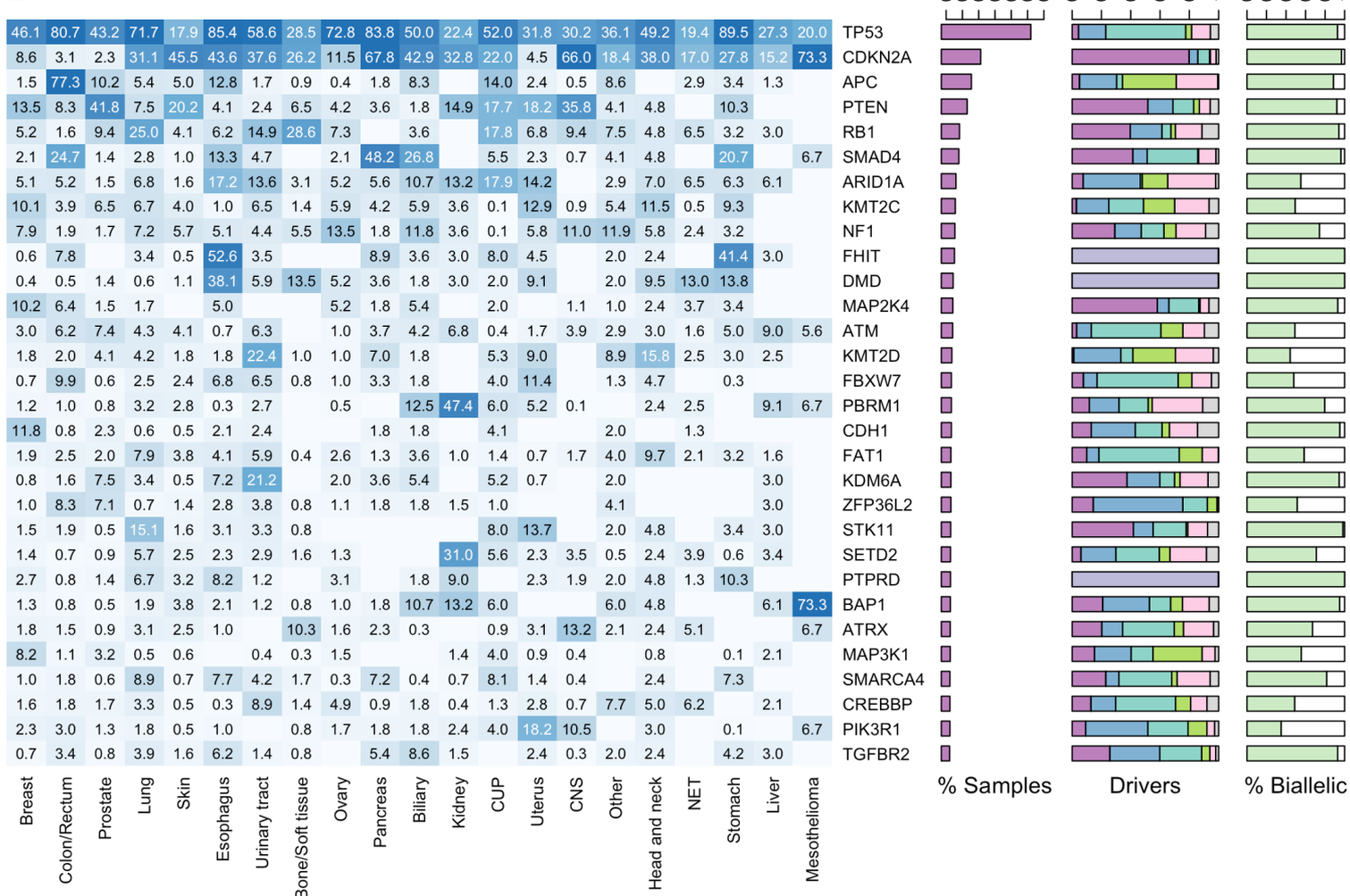
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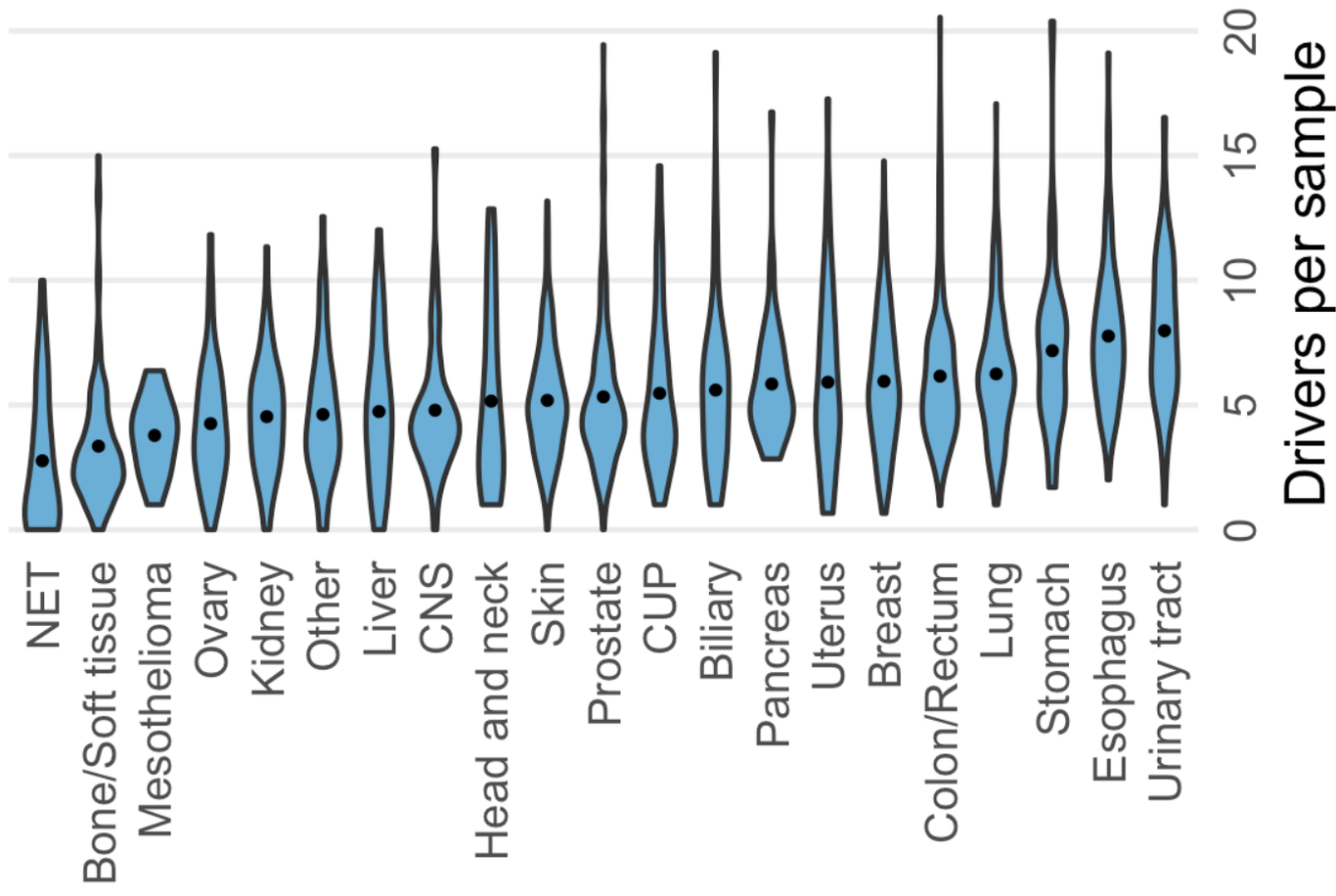
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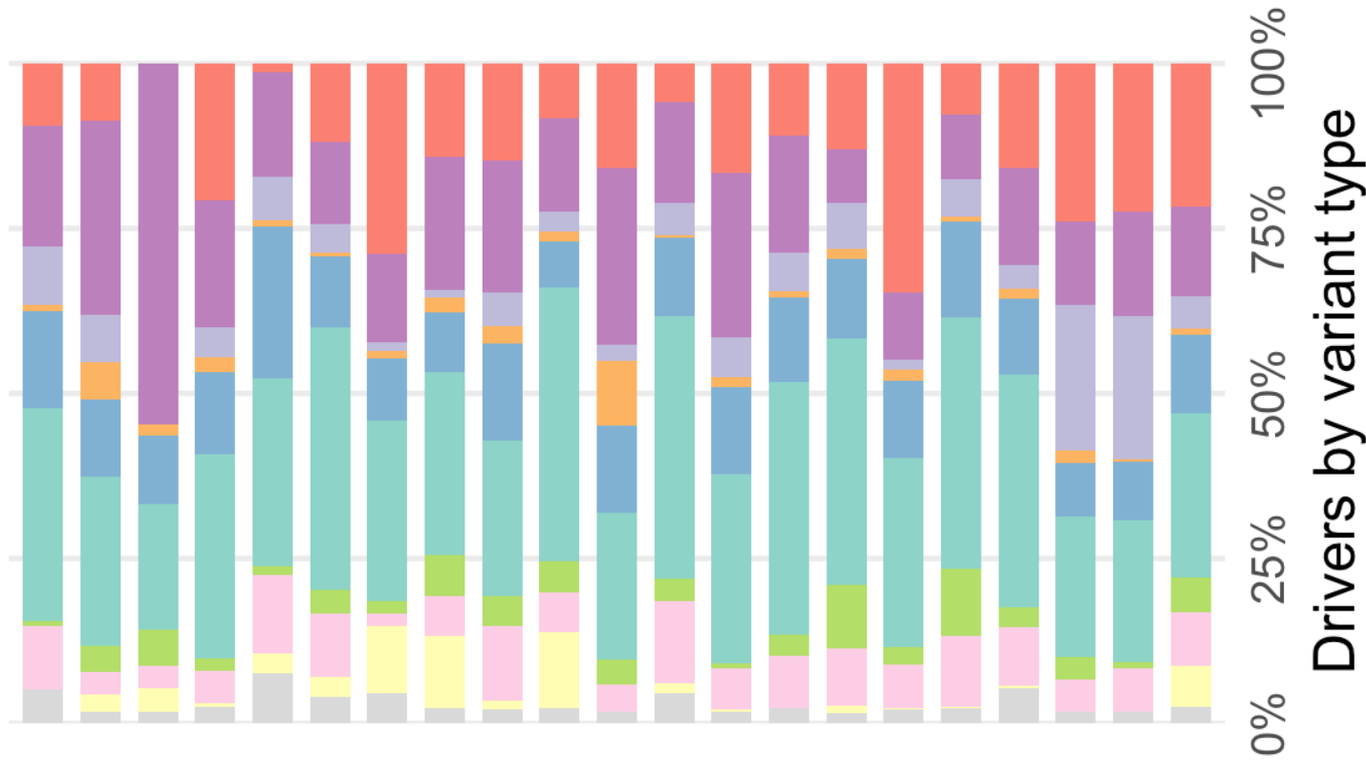
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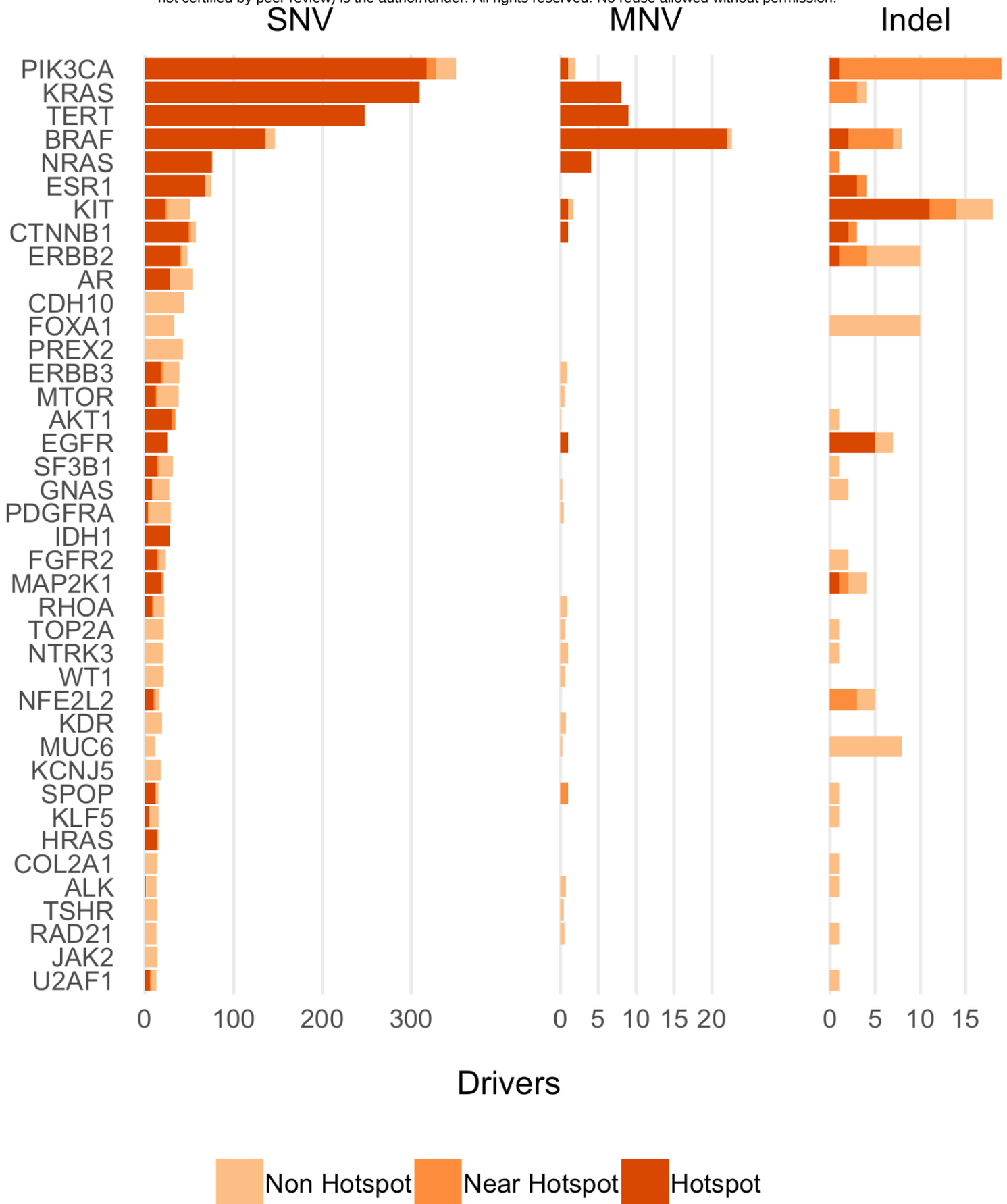
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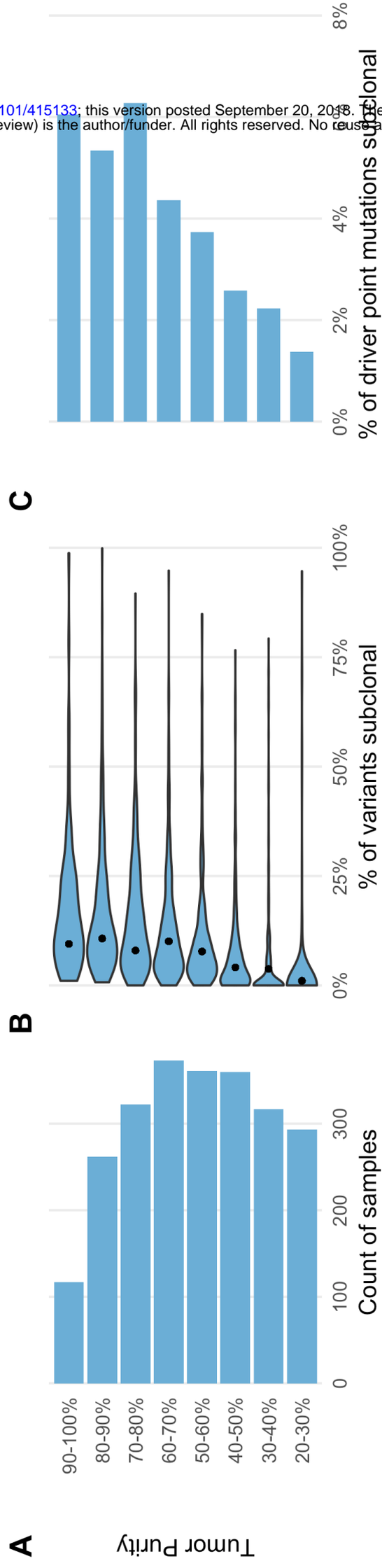


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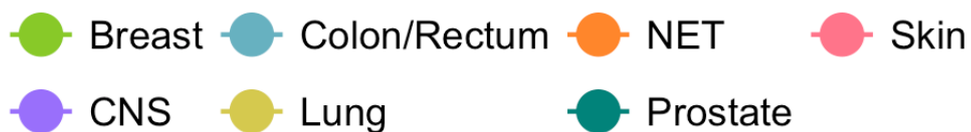
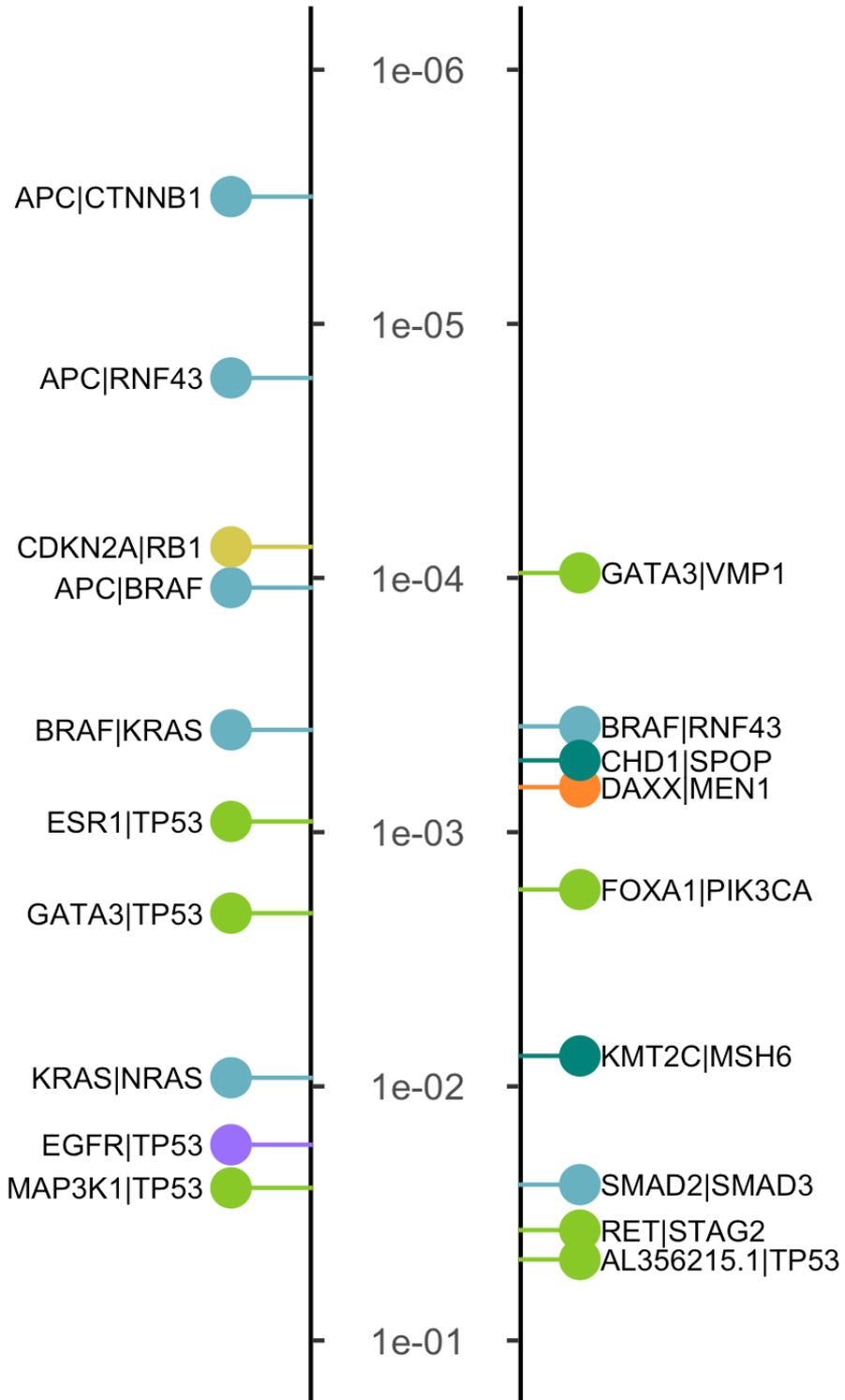
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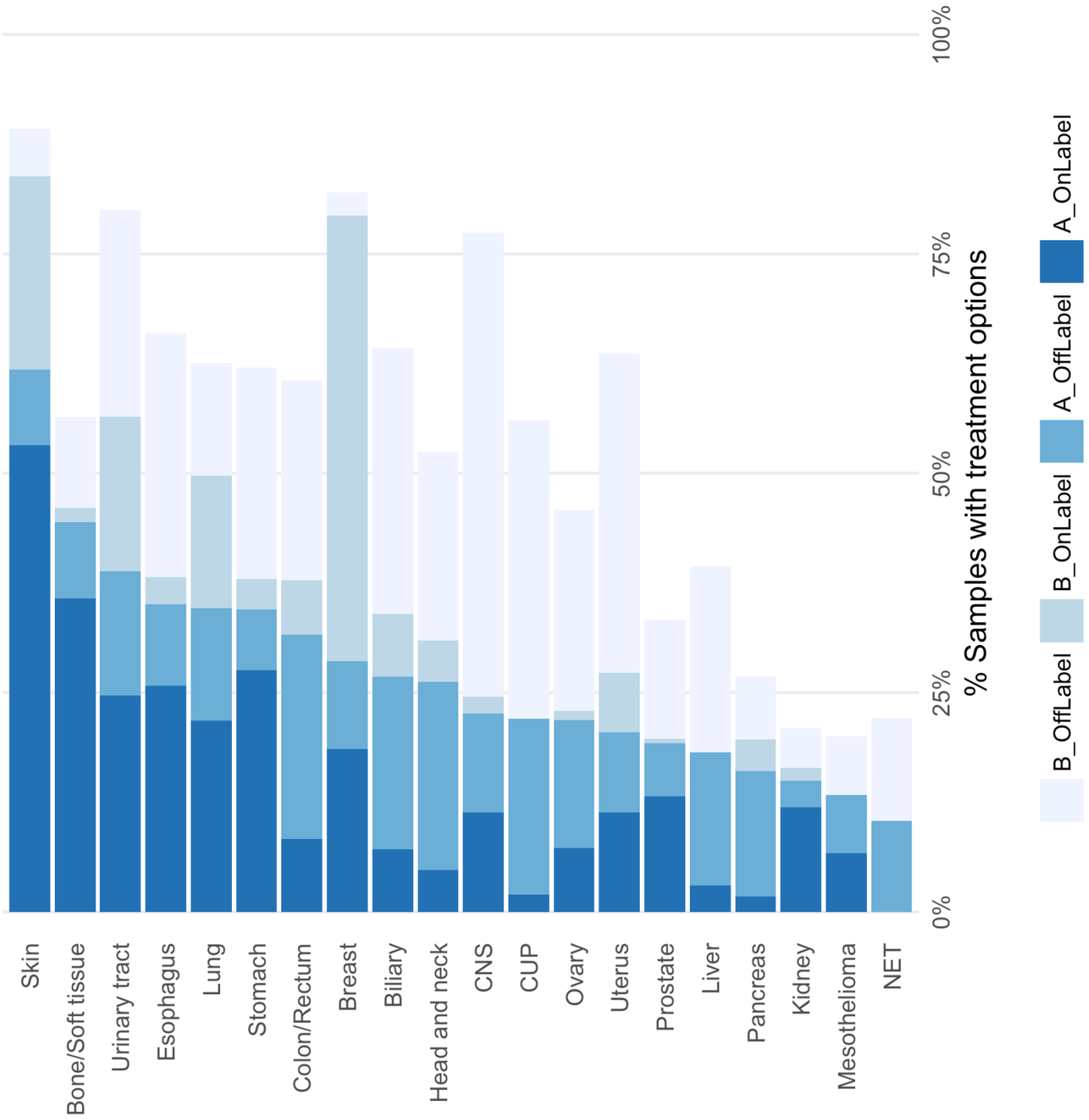
BRAF|NRAS



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Pan-cancer whole genome analyses of metastatic solid tumors

Peter Priestley, Jonathan Baber, Martijn P. Lolkema, Neeltje Steeghs, Ewart de Bruijn, Korneel Duyvesteyn, Susan Haidari, Arne van Hoeck, Wendy Onstenk, Paul Roepman, Charles Shale, Mircea Voda, Haiko J. Bloemendal, Vivianne C.G. Tjan-Heijnen, Carla M.L. van Herpen, Mariette Labots, Petronella O. Witteveen, Egbert F. Smit, Stefan Sleijfer, Emile E. Voest, Edwin Cuppen

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1. Sample collection

Patients with advanced cancer not curable by local treatment options and being candidates for any type of systemic treatment and any line of treatment were included as part of the CPCT-02 (NCT01855477) and DRUP (NCT02925234) clinical studies, which were approved by the medical ethical committees (METC) of the University Medical Center Utrecht and the Netherlands Cancer Institute, respectively. A total of 41 academic, teaching and general hospitals across the Netherlands participated in these studies and collected material and clinical data by standardized protocols¹. Patients have given explicit consent for whole genome sequencing and data sharing for cancer research purposes. Clinical data, including primary tumor type, biopsy location, gender and birth year were collected in electronic case record forms and stored in a central database.

Core needle biopsies were sampled from the metastatic lesion, or when considered not feasible or not safe, from the primary tumor site when still in situ. One to four biopsies were collected (average of 2.1 per patient) and frozen in liquid nitrogen directly after sampling and further processed at a central pathology tissue facility. Frozen biopsies were mounted on a microtome in water droplets for optimal preservation of all types of biomolecules (DNA, RNA and proteins) for subsequent and future omics-based analyses. A single 6 micron section was collected for hematoxylin-eosin (HE) staining and estimation of tumor cellularity by an experienced pathologist. Subsequently, 25 sections of 20 micron, containing an estimated 25,000 to 500,000 cells, were collected in a tube for DNA isolation. In parallel, a tube of blood was collected in CellSave (Menarini-Silicon Biosystems) tubes, which was shipped by room temperature to the central sequencing facility at the Hartwig Medical Foundation. Left-over material (biopsy, DNA) after sample processing was stored in biobanks associated with the studies at the University Medical Center Utrecht and the Netherlands Cancer Institute.

2. Sequencing workflow

DNA was isolated from biopsy and blood on an automated setup (QiaSymphony) according to supplier's protocols (Qiagen) using the DSP DNA Midi kit for blood and QIASymphony DSP DNA Mini kit for tissue and quantified (Qubit). Typically, DNA yield for the tissue biopsy ranged between 50 and 5,000 ng. A total of 50 - 200 ng of DNA was used as input for TruSeq Nano LT library preparation (Illumina), which was performed on an automated liquid handling platform (Beckman Coulter). DNA was sheared using sonication (Covaris) to average fragment lengths of 450 nt. Barcoded libraries were sequenced as pools (blood control 1 lane equivalent, tumor 3 lane equivalents) on HiSeqX (V2.5 reagents) generating 2 x 150 read pairs using standard setting (Illumina).

BCL output from the HiSeqX platform was converted using Illumina bcl2fastq tool (versions 2.17 to 2.20 have been used) using default parameters. Reads were mapped to the reference genome GRCH37 using BWA-mem v0.7.5a², duplicates were marked for filtering and INDELS were realigned using GATK v3.4.46 IndelRealigner. GATK Haplotype Caller v3.4.46 was run to call germline variants in the reference sample. For somatic SNV and INDEL variant calling, GATK BQSR³ is also applied to recalibrate base qualities.

3. Somatic point mutation calling

We called SNV & INDEL somatic variants using Strelka v1.0.14⁴ with the following optimisations:

- **Preservation of known variants:** From the raw Strelka output we marked all known pathogenic variants from external databases such that these would be preserved from all subsequent filtering. The list of pathogenic variants used was the union of:

- Point mutations in CIViC⁵ with level C evidence or higher (download = 01-mar-2018)
- Somatic variants from CGI⁶ (update: 17-jan-2018)
- Oncogenic or likely Oncogenic variants from OncoKb⁷ (download = 01-mar-2018); <http://oncokb.org/api/v1/utills/allAnnotatedVariants.txt>
- TERT promoter variants at genomic coordinates: 5:1295242, 5:1295228, 5:1295250
- **Modified quality score filtering**
 - We split variants into high confidence (HC) and low confidence (LC) regions using the NA12878 GIABv3.2.2 high confidence region definitions⁸, based on the observation that we produce far higher rates of false positives variant calls in LC regions
 - Set quality score cutoffs for SNV & INDEL to 10 for HC regions and 20 for LC regions (default = 15 for SNV, 30 for INDEL)
 - Added an additional quality filter to tighten filtering for low allelic frequency variants: quality score * allele frequency > 1.3
- **Improved repeat sensitivity:** Switched off the default Strelka repeat filter to improve indel calling in microsatellites and short repeats.
- **Panel of normals (PON) to remove germline leakage:** Filtered out any variants which were found by GATK haplotypcaller in more than 5 samples in a germline PON consisting of 2000 of our reference blood samples. PON available at (<https://resources.hartwigmedicalfoundation.nl/>)
- **PON to remove strelka-specific artefacts:** Filtered any variant which was supported by 2 or more reads in strelka in the reference sample in at least 4 patients in our cohort. PON available at (<https://resources.hartwigmedicalfoundation.nl/>)
- **Removal of INDELS near a PON filtered INDEL -** Regions of complex haplotype alterations are often called as multiple long indels which can make it more difficult to construct an effective PON, and sometimes we find residual artefacts at these locations. Hence we also filter inserts or deletes which are 3 bases or longer where there is a PON filtered INDEL of 3 bases or longer within 10 bases in the same sample.

The settings and tools for this optimized HMF pipeline are available at <https://github.com/hartwigmedical/>. We tested the default and HMF optimized settings on a GIAB mix-in sample (ref = NA24385, tumor = 70% NA24385, 30% NA12878) to test sensitivity at a realistic purity and on a null tumor (ref = NA12878, tumor = NA12878) to test precision. The results of this analysis are as follows:

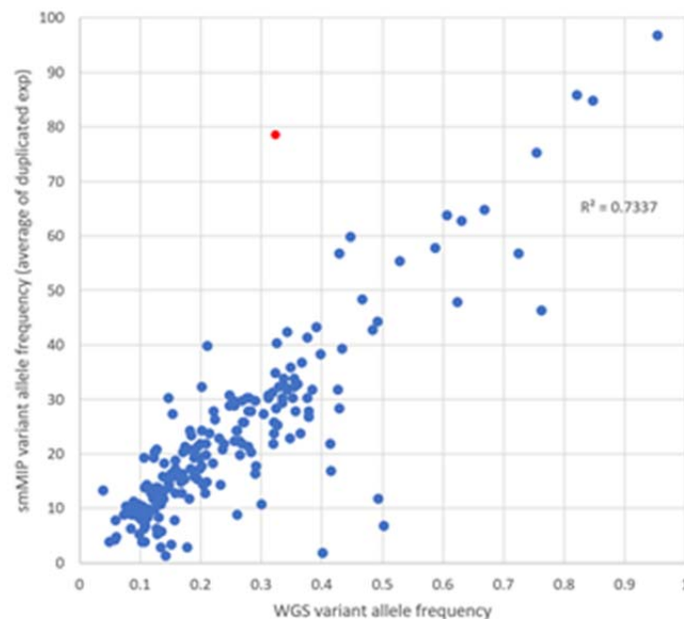
Configuration	SNV sensitivity	SNV false positive / genome	INDEL sensitivity	Indel false positive / genome
Default	93%	3500	24%	41
Optimized HMF pipeline	96%	109	77%	27

We performed external validation of a set of single nucleotide variants (SNV) and short insertion/deletions (indels) that have been detected by Whole Genome Sequencing (WGS) using the single molecule Molecular Inversion Probe (smMIP) technology⁹. SNV and short indels variants were semi-randomly selected from 30 patient samples. The first selection was to include every variant that was reported in the patient report (114 gene panel) as obtained during the routine CPCT-02 study analysis. This way, a total of 82 variants (67 SNVs, 15 indels) were selected in 45 genes. The second selection involved random sampling of 256 coding and non-coding variants from the same 30 patient samples.

A custom smMIP panel was designed to cover the selected variants. For 45 variants (17.6%) no smMIP design was possible, all of which were intergenic variants. For the other 211 variants probes could successfully be designed. Analysis of the smMIP sequencing data indicated that for 17 of the 211 variants

(8.1%) the smMIP sequencing data was of insufficient quality (mostly due to repeat stretches), while the WGS data seemed sufficiently reliable for accurate calling (confirmed by visual inspection of the read data), including 3 coding variants (*RB1*, *ERBB4* and *BRCA2*) and 14 intergenic regions. The retrospective investigation of the WGS data indicated that for another three variants (1.4%) the smMIP as well as the WGS data was of insufficient quality due to large homopolymer stretches.

In total 192 variants could be successfully sequenced and analyzed using the smMIP and could be used for confirmation of the WGS findings. 189 SNVs and indel variants (98.4%) were confirmed by smMIP sequencing, indicating a very high accuracy of WGS-derived variant calling results. All three variants that could not be confirmed by smMIP were from intergenic regions, including 1 variant that showed a mixed double-variant (chr3:75887550_G>T/C) and for which both technologies had difficulties in accurately calling the genotype. For the remaining 2 variants (chr8:106533360_106533361insAC, chr12:125662751_125662752insA), it remains unclear if these could not be detected by smMIP or were falsely called by WGS, as they fall in repetitive genomic stretches.



The 189 successfully confirmed variants showed a good linear correlation in variant allele frequency between WGS and smMIP sequencing (average of duplicates) with an R^2 of 0.733. This result indicated that WGS, with its lower read depth (on average between 100-110x) than smMIP and without a read-barcoding system, is accurate in quantitatively determining the variant frequency at frequencies above 5%. One variant (chr19:55276095C>T, indicated in red in the figure above) showed a large deviation in variant frequency, which was likely caused due to the much lower than expected coverage of the variant, both in the WGS (37 reads) as well as in the smMIP data (28 and 35 reads).

4. MNV correction

Strelka somatic variants that appear on consecutive positions, or 1 base apart were considered potential multi nucleotide variants (MNVs). The BAM files were re-examined, and the variants were merged into a single MNV if greater than 80% of the reads with a mapping quality score of at least 10 and which are neither unmapped, duplicated, secondary, nor supplementary containing any of the individual variants also contained the other variants of the potential MNV. The attributes of the resulting MNV

variant were determined by picking the minimum values from the individual variants forming the MNV. MNVs were marked as PON filtered only if both individual variants were PON filtered.

5. Somatic structural variant calling

Structural Variants were called using Manta(v1.0.3)¹⁰ with default parameters. We then re-examined each breakpoint calculated variant allele frequencies for each break end and applied seven additional filters to the Manta output to improve precision using an internally built tool called 'Breakpoint-Inspector' (BPI) v1.5. Two main types of filters are applied by BPI:

- **Evidence of variant in reference sample** - variants are filtered if we can find any evidence of paired read support, split read support or soft clipping concordance (5+ bases at exact breakpoint) in the matching blood sample.
- **Inadequate support for variant in tumor sample** - For all inversions and translocations and for long deletions and tandem duplications (>1000 bases between breakpoints) we require at least 1 read with paired read support. For short deletions and duplications (<1000 bases between breakpoints) we require at least 1 read with split read support. In both cases at least one of those reads must be anchored with at least 30 bases at each breakpoint. We also require the minimum read coverage across each breakpoint in the tumor to be > 10 depth.

Code and description of filters for BPI are available at <https://github.com/hartwigmedical/hmftools/tree/master/break-point-inspector>.

Each break end was annotated with its position in all transcripts from 'KNOWN' genes in Ensembl v89.37¹¹. Each gene was marked as disrupted if there was at least one structural variant that impacted on the canonical transcript.

6. Identification of gene fusions

For each structural variant, every combination of annotated overlapping transcripts from each breakend was tested to see if it could potentially form an intronic inframe fusion. A list of 411 curated known fusion pairs was sourced by taking the union of known fusions from the following external databases:

- Cosmic curated fusions¹² (v83)
- OncoKb⁷ (download = 01-mar-2018)
- CGI⁶ (update: 17-jan-2018)
- CIViC⁵ (download = 01-mar-2018)

We then also created a list of promiscuous fusion partners using the following rules

- **3' promiscuous:** Any gene which appears on the 3' side in more than 3 of the curated fusion pairs OR appears at least once on the 3' side and is marked as promiscuous in either OncoKb, CGI or CIViC
- **5' promiscuous:** Any gene which appears on the 5' side in more than 3 of the curated fusion pairs OR appears at least once on the 5' side and is marked as promiscuous in either OncoKb, CGI or CIViC

For each promiscuous partner we also curated a list of essential domains that must be preserved to form a viable fusion partner.

Finally, we report an intronic inframe fusion if the following conditions are met

- Matches an exact fusion from the curated list OR is intergenic and matches 5' promiscuous OR matches 3' promiscuous gene

- Curated domains are preserved
- Does not involve the 3'UTR region of either gene
- For intragenic fusions, must start and end in coding regions of the gene

7. Copy number calling

We use an in house developed integrated tool, PURity & PLOidy Estimator (PURPLE), that combines B-allele frequency (BAF), read depth and structural variants to estimate the purity and copy number profile of a tumor sample. Version v2.14 of PURPLE has been used.

There are 5 key steps in the PURPLE pipeline:

1. Calculate BAF in tumor at high confidence heterozygous germline loci

We determine the BAF of the tumor sample by finding heterozygous locations in the reference sample from a panel of 796,447 common germline heterozygous SNP locations. To ensure that we only capture heterozygous points, we filter the panel to only loci with allelic frequencies in the reference sample between 40% and 60% and with depth between 50% and 150% of the reference sample genome wide average. Typically, this yields 140k-200k heterozygous germline variants per patient. We then calculate the allelic frequency of corresponding locations in the tumor.

2. Determine read depth ratios for tumor and reference genomes

The raw read counts per 1,000 base window for both normal and tumor samples, by counting the number of alignment starts in the respective bam files with a mapping quality score of at least 10 that is neither unmapped, duplicated, secondary, nor supplementary. Windows with a GC content less than 0.2 or greater than 0.6 or with an average mappability below 0.85 are excluded from further analysis.

Next we apply a GC normalization to calculate the read ratios. We divide the read count of each window by the median read count of all windows sharing the same GC content then normalise further to the ratio of the median to mean read count of all windows.

Finally, the reference sample ratios have a further 'diploid' normalization applied to them to remove megabase scale GC biases. This normalization assumes that the median ratio of each 10Mb window (minimum 1Mb readable) should be diploid for autosomes and haploid for sex chromosomes in males in the germline sample.

3. Segmentation

We segment the genome into regions of uniform copy number by combining segments generated from the read ratios for both tumor and reference sample, from the BAF points with structural variant breakpoints derived from Manta & BPI. Read ratios and BAF points are segmented independently using the Bioconductor copynumber package¹³ which uses a piecewise constant fit (PCF) algorithm (with custom settings gamma = 100, k = 1). These segment breaks are then combined with the structural variants breaks according to the following rules:

1. Every structural variant break starts a new segment, as does chromosome starts, ends and centromeres. This is regardless of if they are distinguishable from existing segments or not.
2. Ratio and BAF segment breaks are only included if they are distinguishable from an existing segment.
3. To be distinguishable, a break must be at least one complete mappable read depth window away from an existing segment.

Once the segments have been established we map our observations to them. In each segment we take the median BAF of the tumor sample and the median read ratio of the tumor and reference samples. We also record the number of BAF points within the segment as the BAFCount.

A reference sample copy number status is determined at this stage based on the observed copy number ratio in the reference sample, either 'DIPLOID' (0.8<= read depth ratio<=1.2), 'HETEROZYGOUS_DELETION' (0.1<=ratio<0.8), 'HOMOZYGOUS_DELETION' (ratio<0.1), 'AMPLIFICATION'(1.2<ratio<=2.2) or 'NOISE' (ratio>2.2). The purity fitting and smoothing steps below use only the DIPLOID germline segments.

4. Purity Fitting

Next we jointly fit tumor purity and sample ploidy (expressed as a normalisation factor) according to the following principles:

1. The absolute copy number of each segment should be close to an integer ploidy
2. The BAF of each segment should be close to a % implied by integer major and minor allele ploidies.
3. Higher ploidies have more degenerate fits but are less biologically plausible and should be penalised
4. Segments are weighted by the count of BAF observations which is treated as a proxy for confidence of BAF and read depth ratio inputs.
5. Segments with lower observed BAFs have more degenerate fits and are weighted less in the fit

For any given tumor purity and sample ploidy we calculate the score by first modelling the major and minor allele ploidy of each segment and minimising the deviation between the observed and modelled values according to the following formulas:

$$\begin{aligned} \text{ModelDeviation} &= \text{abs}(\text{ObservedRatio} - \text{ModelRatio}) + \text{abs}(\text{ObservedBaf} - \text{ModelBaf}) \\ \text{ModelBaf} &= (\text{tumorPurity} * (\text{segmentMinorPloidy} - 1) + 1) / (\text{tumorPurity} * (\text{segmentPloidy} - 2) + 2) \\ \text{ModelRatio} &= \text{sampleNormFactor} + (\text{segmentPloidy} - 2) * \text{tumorPurity} * \text{sampleNormFactor} / 2d; \end{aligned}$$

Once modelled, each segment is given a ploidy penalty:

$$\begin{aligned} \text{PloidyPenalty} &= 1 + \min(\text{SingleEventDistance}, \text{WholeGenomeDoublingDistance}); \\ \text{WholeGenomeDoublingDistance} &= 1 + \text{abs}(\text{segmentMajorAllele} - 2) + \text{abs}(\text{segmentMinorAllele} - 2); \\ \text{SingleEventDistance} &= \text{abs}(\text{segmentMajorAllele} - 1) + \text{abs}(\text{segmentMinorAllele} - 1); \end{aligned}$$

Summing up over all the segments generates a score for each tumor purity / sample ploidy combination from which we can select the minimum:

$$\begin{aligned} \text{FittedPurityScore} &= \frac{1}{\text{TotalBafCount}} \sum_{i=1}^n \text{PloidyPenalty}_i \times \text{ModelDeviation}_i \times \text{BafCount}_i \\ &\times \text{ObservedBaf}_i \end{aligned}$$

Given a fitted purity and sample ploidy we are then able to determine the purity adjusted copy number and BAF of each segment in the tumor from the unadjusted read ratios and BAFs respectively.

The purity estimates of PURPLE were validated using the tumor cell line COLO829. We created diluted in-silico mixture models of the tumor and blood cell lines for COLO829 with simulated purities of 20%, 30%, 40%, 60%, 80% and 100%, and ran PURPLE on the simulated BAM files against a reference sample.

The PURPLE estimates match the simulation closely as follows:

Simulated Purity	PURPLE estimated purity	Difference
20%	20%	0%
30%	30%	0%
40%	40%	0%
50%	50%	0%
60%	60%	0%
80%	81%	1%
100%	100%	0%

5. Smoothing

Since the segmentation algorithm is highly sensitive, and there is a significant amount of noise in the read depth in whole genome sequencing, many adjacent segments created above will have a similar copy number and BAF profile and can be combined and averaged to form a larger, smoothed, region.

We apply a number of rules to merge adjacent regions to create a smooth copy number profile.

1. Never merge a segment break created from a structural variant break end.
2. Use the count of BAF points as a proxy for confidence or weight in the region. Note that some segments may have a BAF count of 0.
3. Merge segments where the difference in BAF and copy number is within tolerances.
4. BAF tolerance is linear between 0.03 and 0.35 dependent on BAF count.
5. Copy number tolerance is linear between 0.3 and 0.7 dependent on BAF count. The tolerance also increases linearly as purity of the tumor sample decreases below 20%.
6. Start from most confident segment and smooth outwards until we reach a segment outside of tolerance. Move on to next highest unsmoothed section.
7. It is possible to merge in (multiple) segments that would otherwise be outside of tolerances if:
 - a. The total dubious region is sufficiently small (<30k bases or <50k bases if approaching centromere); and
 - b. The dubious region does not end because of a structural variant; and
 - c. The dubious region ends at a centromere, telomere or a segment that is within tolerances.
8. When the entire short arm of a chromosome is lacking copy number information (generally on chromosome 13,14,15,21, or 22), the copy number of the long arm is extended to the short arm.
9. Any remaining unknown segments are given the expected copy number of their associated chromosome, i.e. 2 for autosomes and female allosomes, 1 for male allosomes.

Where clusters of SVs exist which are closer together than our read depth ratio window resolution of 1,000 bases, the segments in between will not have any copy number information associated with them.

To resolve this, we infer the ploidy from the surrounding copy number regions. The outermost segment of any SV cluster will be associated with a structural variant with a ploidy that can be determined from the adjacent copy number region and the VAF of the SV. We use this ploidy and the orientation of structural variant to calculate the change in copy number across the SV and hence the copy number of the outermost unknown segment. We repeat this process iteratively and infer the copy number of all regions within a cluster.

Once region smoothing is complete, it is possible there will be regions of unknown BAF, if no BAF points were present in a copy number region. We infer this BAF by assuming that they share their minor allele ploidy with their neighbouring region. If there are multiple neighbouring regions with known BAF we use the highest confident region (i.e. highest BAF count) to infer.

At this stage we have determined a copy number and minor allele ploidy for every base in the genome.

8. Sample filtering based on copy number output

Following our copy number calling, samples were QC filtered from the analysis based on 4 criteria:

- **NO_TUMOR** - If PURPLE fails to find any aneuploidy AND the number of somatic SNVs found is less than 1,000 then the sample is marked as NO_TUMOR.
- **MIN_PURITY** - We exclude samples with a purity of <20%
- **FAIL_SEGMENT** - We remove samples with more than 120 copy number segments unsupported at either end by SV breakpoints. This step was added to remove samples with extreme GC bias, with differences in depth of up to or in excess of 10x between high and low GC regions. GC normalisation is unreliable when the corrections are so extreme so we filter.
- **FAIL_DELETED_GENES** - We removed any samples with more than 280 deleted genes. This QC step was added after observing that in a handful of samples with high MB scale positive GC bias we sometimes systematically underestimate the copy number in high GC regions. This can lead us to incorrectly infer homozygous loss of entire chromosomes, particularly on chromosome 17 and 19.

Where multiple biopsies exist for a single patient, we always choose the highest purity sample for our analysis of mutational load and drivers.

9. Assessment of impact of sequencing depth coverage on variant calling sensitivity

To assess the impact of our sequencing depth on variant calling sensitivity, we selected 10 samples at random, downsampled the BAMs by 50%. We then reran the identical somatic variant calling pipeline.

Comparing the output to the original runs, we found near identical purities and ploidies for the down sampled runs ([Extended Data Fig. 2](#)). We observed an average decrease in sensitivity of 10% for SNV, 15% for MNV, 19% for SV, and 2% for INDEL.

10. Clonality and biallelic status of point mutations

For each point mutation we determined the clonality and biallelic status by comparing the estimated ploidy of the variant to the local copy number at the exact base of the variant. The ploidy of each variant is calculated by adjusting the observed VAF by the purity and then multiplying by the local copy number to work out the absolute number of chromatids that contain the variant.

We mark a mutation as biallelic (i.e. no wild type remaining) if Variant Ploidy > Local Copy Number - 0.5. The 0.5 tolerance is used to allow for the binomial distribution of VAF measurements for each variant. For example, if the local copy number is 2 than any somatic variant with measured ploidy > 1.5 is marked as biallelic.

For each variant we also determine a probability that it is subclonal. This is achieved via a 2 step process

1. Fit the somatic ploidies for each sample into a set of clonal and subclonal peaks

We apply an iterative algorithm to find peaks in the ploidy distribution:

- Determine the peak by finding the highest density of variants within +/- 0.1 of every 0.01 ploidy bucket.
- Sample the variants within a 0.05 ploidy range around the peak.
- For each sampled variant, use a binomial distribution to estimate the likelihood that the variant would appear in all other 0.05 ploidy buckets.
- Sum the expected variants from the peak across all ploidy buckets and subtract from the distribution.
- Repeat the process with the next peak

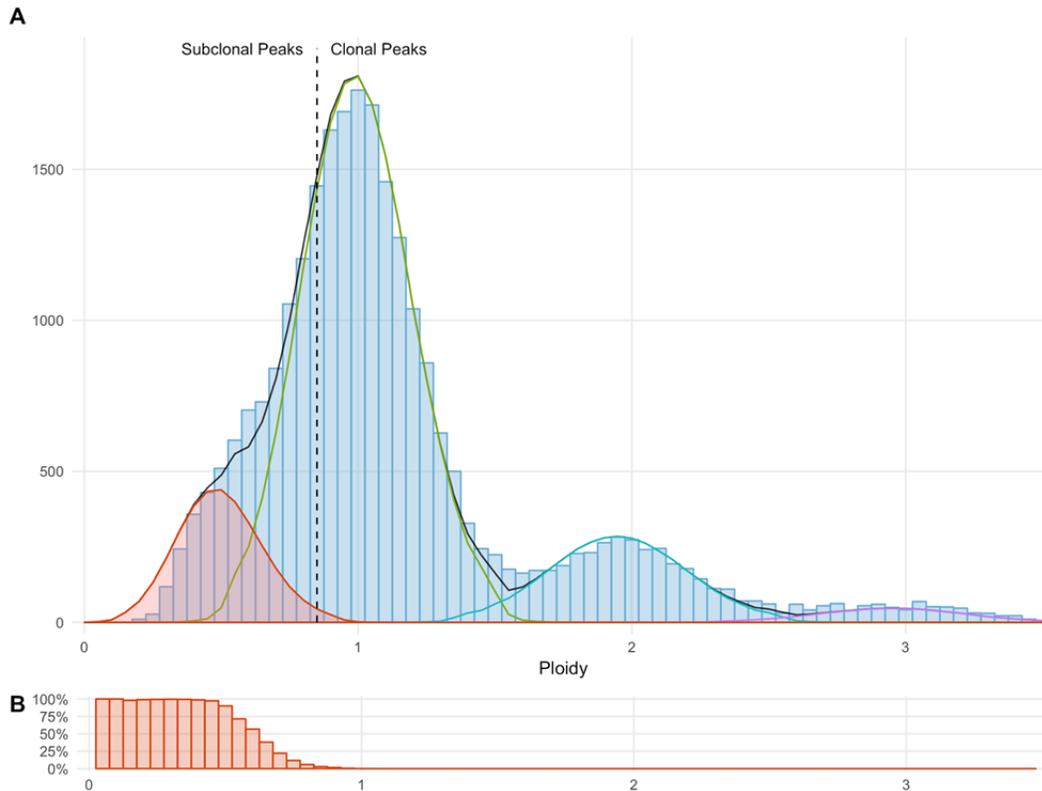
This process yields a set of ploidy peaks, each with a ploidy and a total density (i.e. count of variants). To avoid overfitting small amounts of noise in the distribution, we filter out any peaks that account for less than 40% of the variants in the ploidy bucket at the peak itself. After this filtering we scale the fitted peaks by a constant so that the sum of fitted peaks = the total variant count of the sample.

Finally we mark a peak as subclonal if the peak ploidy < 0.85

2. Calculate the probability that each individual variant belongs to each peak

Once we have fitted the somatic ploidy peaks and determined their clonality, we can calculate the subclonal likelihood for any individual variant as the proportion of subclonal variants at that same ploidy.

The following diagram illustrates this process for a typical sample. Figure A shows the histogram of somatic ploidy for all SNV and INDEL in blue. Superimposed are four peaks in different colours fitted from the sample as described above. The red filled peak is below the 0.85 threshold and is thus considered subclonal. The black line shows the overall fitted ploidy distribution. Figure B shows the likelihood of a variant being subclonal at any given ploidy.



Subclonal counts in this paper are calculated as the total density of the subclonal peaks for each sample. Subclonal driver counts are calculated as the sum across the driver catalog of subclonal probability * driver likelihood (driver likelihood is explained in detail below).

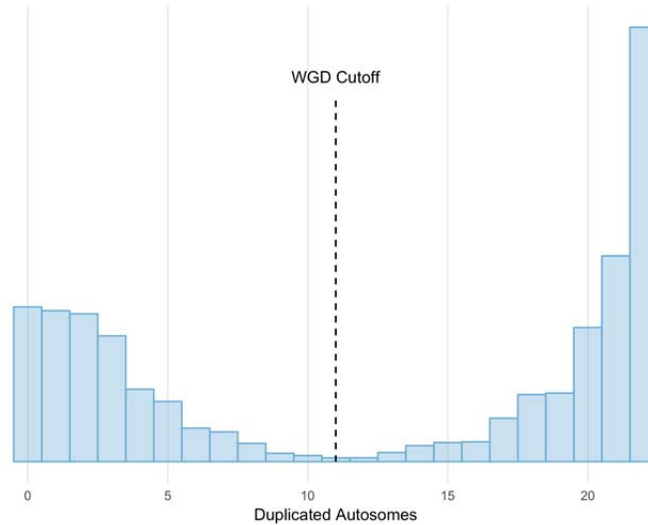
11. WGD status determination

We implement a simple heuristic that determines if Whole Genome Duplication has occurred:

Major allele Ploidy >1.5 on at least 50% of at least 11 autosomes

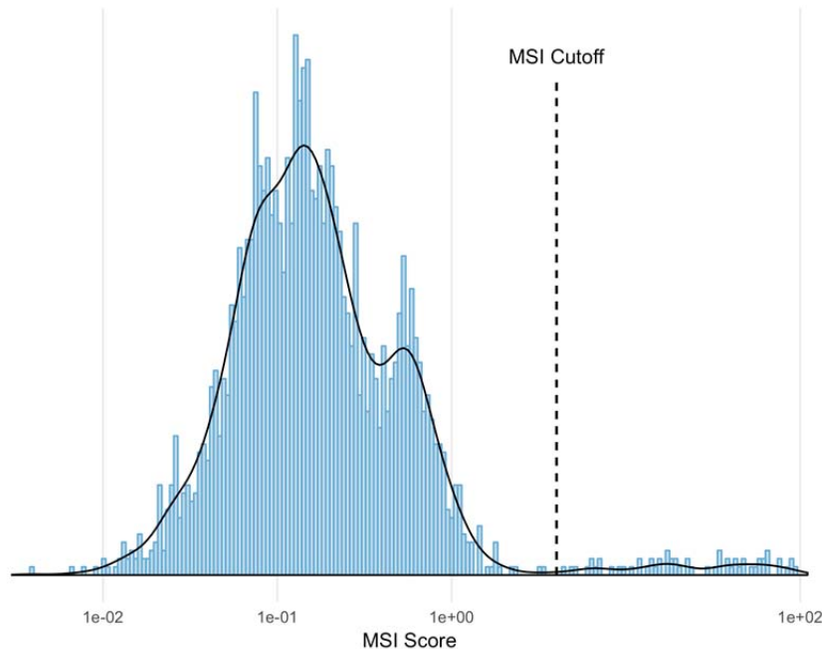
The principle behind this heuristic is that if sufficient independent chromosomes are predominantly duplicated, the most parsimonious explanation is that the duplication occurred in a single genome-wide event.

The number of duplicated autosomes per sample (ie. the number of autosomes which satisfy the above rule) follows a bimodal distribution with 95% of samples have either ≤ 6 or ≥ 15 autosomes duplicated. Hence, the classification of a genome as WGD is not particularly sensitive to the choice of cut-off as is evident the following chart:



12. MSI status determination

To determine the MSI status of all samples we used the method described by the MSIsSeq tool¹⁴. In brief, we count the number of INDELS per million bases occurring in homopolymers of 5 or more bases or dinucleotide, trinucleotide and tetranucleotide sequences of repeat count 4 or more. MSIsSeq scores ranged from 0.004 up to 98.63, with a long tail towards lower MSI scores as shown in the following chart:



To be able to accurately set and validate the MSIsSeq cutoff for classification of MSI we compared the WGS results with the standard, routinely used MSI assessment using a 5-marker PCR panel (BAT25, BAT26, NR21, NR24 and MONO27 markers). For a batch of 48 pre-selected samples, the MSI PCR assay was blindly performed by an independent ISO-accredited pathology laboratory. Both the binary MSI and MSS classifications were determined, but also the number of positive markers.

A sample was considered as MSI if two or more of the five markers were score as positive (instable). PCR-based analysis identified 16 MSI samples, all of which were also identified by MSIseq with scores >4. MSIseq identified one sample that was missed by PCR-based analysis, although this sample showed microsatellite instability for one out the five markers. The MSIseq scores thus highly correlate with the number of positive MSI PCR markers and all, except one, samples with an elevated score are classified as MSI by pathology. Based on this data we determined the best cutoff for MSIseq classification to be at a **score of 4**.

Results of the PCR-based and WGS based MSI classification are summarized in the table below. The sensitivity of WGS-based MSI classification on this set was 100% (95%CI 82.6 – 100%) with a specificity of 97% (95%CI 88.2-96.9%). The calculated Cohen's kappa score was 0.954 (95%CI 0.696-0.954), indicative of a very high agreement.

	PCR-MSS	PCR-MSI	Total
MSIseq -MSS	31	0	31
MSIseq - MSI	1	16	17
Total	32	16	48

13. Holistic gene panel for driver discovery

We used Ensembl release 89 as a basis for our gene definitions and have taken the union of Entrez identifiable genes and protein coding genes as our base panel.

Certain genes have multiple definitions. NPIPA7 for example has two definitions which are equally valid, [ENSG00000214967](#) and [ENSG00000183889](#). To solve this we select a single gene definition based on the following steps:

- 1) Exclude non protein coding genes.
- 2) Favour genes that are present in both Havana and Ensembl.
- 3) Select gene with longest transcript.

This returns our final gene panel tally to 25,963 genes of which 20,083 genes are protein coding. For each gene we chose the canonical transcript or the longest if no canonical transcript exists.

For CDKN2A, we included both the p16 and p14arf transcripts in the analysis given the known importance of both transcripts to tumorigenesis¹⁵ and the fact that the two transcripts use alternate reading frames in the same exon.

14. Significantly mutated driver genes discovery

Using all SNV and INDEL variants from the holistic gene panel, we ran dNdScv¹⁶ to find significantly mutated genes (SMGs) and also to estimate the proportion of missense, nonsense, essential splice site and INDEL variants which are drivers in each individual gene in the panel.

Pan cancer and at an individual cancer level we tested the normalised dNdS rates against a null hypothesis that dNdS = 1 for each variant subtype. To identify SMGs in our cohort we used a strict significance cutoff of $q < 0.01$.

2 of the HMF SMG candidates were subsequently removed via manual curation as they were deemed to be likely artefacts of our methods:

- POM121L12 - found only to be significant due to an extreme covariate value in dndscv
- TRIM49B - found to have poor mappability on nearly all its variants and a known close paralog

15. Significantly amplified & deleted driver gene discovery

To search for significantly amplified and deleted genes we first calculated the minimum exonic copy number per gene across our holistic gene panel. For amplifications, we searched for all the genes with high level amplifications only (defined as minimum Exonic Copy number $> 3 * \text{sample ploidy}$). For deletions, we searched for all the genes in each sample with either full or partial gene homozygous deletions (defined as minimum exonic copy number < 0.5). The Y chromosome was excluded from the deletion analysis since the Y chromosome is deleted altogether in 35% of all male cancer samples in our cohort and hence is difficult to distinguish at the gene level.

We then searched separately for amplifications and deletions, on a per chromosome basis, for the most significant focal peaks, using an iterative GISTIC-like peel off method¹⁷, specifically:

- Find the highest scoring gene.
 - For deletions the score is simply the count of samples with homozygous deletions in the gene.
 - For amplifications, we need to consider both the count and strength of the amplification so we use:
 - $\text{score} = \text{sum}(\log_2(\text{copy number} / \text{sample ploidy}))$.
- Record gene as a peak, and mark all consecutive genes with a score within 15% and 25% of the highest score for deletions and amplifications respectively as part of the candidate peak.
- 'Peel' off all samples which contributed to the peak across the entire chromosome
- Repeat the process

A filter was applied where we removed deletions from a handful of noisy copy number regions in the genome where we found more than 50% of the observed deletions were not supported on either breakend by a structural variant.

Most of the deletion peaks resolve clearly to a single target gene reflecting the fact that homozygous deletions are highly focal, but for amplifications this is not the case and the majority of our peaks have 10 or more candidates. We therefore annotated the peaks, to choose a single putative target gene using an objective set of automated curation rules in order of precedence:

- If more than 50% of the copy number events in the peeled samples include the telomere or centromere then mark as $\langle \text{CHR} \rangle _ \langle \text{ARM} \rangle _ \langle \text{TELOMERE/CENTROMERE} \rangle$
- Else choose highest scoring candidate gene which matches a list of actionable amplifications from OncoKB, CGI and CIViC clinical annotation DBs.
- Else choose highest scoring candidate gene found in our panel of significantly mutated genes
- Else choose highest scoring candidate gene found in cosmic census
- Else choose highest scoring protein coding candidate gene
- Else choose longest non-coding candidate gene

Finally, we filter the peaks to only highly significant deletions and amplifications using the following rules

- Deletions => Keep any peak with > 5 homozygous deletions
- Amplifications => Keep any peak with score > 29

These cut-offs were chosen using a binomial model which assumes the probability of any given gene being observed to be randomly deleted or highly amplified is equal to the average number of genes amplified or deleted in each event divided by the total number of genes considered. The cut-offs were chosen to be the lowest score with a q-value below 0.25. Since amplifications are generally much broader

(averaged genes affected per event of 41.6 compared to just 5.4 for deletions) a much higher number of genes is required to reach significance.

The calculation details for the cut-offs are presented in the table below.

	Cohort Data						Statistical Calculations				
	Count of Events	Sum Scores	Count of genes affected	Avg Genes affected per event	Avg score / event	Total Genes Tested	Probability event overlaps a given gene	Score cutoff	P Value of cutoff	Significant findings	Q Value
Dels	4,915	4,915	26,676	5.4	1.0	25,965	0.00021	5	0.00068	117	0.15
Amps	3,925	6,959	163,393	41.6	1.8	25,965	0.00160	29	0.00030	33	0.23

This model is likely to be highly conservative as it assumes that all the events are passengers, whereas in fact a high proportion contain driver genes.

16. Fragile site annotation

Homozygous deletions were also annotated as common fragile site (CFS) based on their genomic characteristics. This annotation is not definitive, but is useful as CFS are known to be regions of high genomic instability. Hence despite being significantly deleted, their status as a genuine cancer driver remains unclear.

There is no absolute agreement on which regions should be classified as CFS, but two well-known features are a strong enrichment in long genes and a high rate of observed deletions of up to 1 megabase¹⁸. Hence for this analysis we classified a gene as a fragile site if it met all the following criteria:

- Total length of gene > 500,000 bases
- More than 30% of all SVs with breakpoints that disrupt the gene are deletions with length greater than 20,000 bases and less than 1 megabase.
- The gene is not found to be significantly mutated (by dNdScv) in our cohort or in Martincorena et al.¹⁶

Using these criteria we annotated the following list of 16 Genes as fragile:

Gene	Chr	Start position	Length (bases)	Total Disruptive SV Count	% of SV that are DELs (>20kb & <1MB)
LRP1B	2	140,988,992	1,900,278	1,272	0.469
FHIT	3	59,735,036	1,502,097	2,128	0.596
LSAMP	3	115,521,235	2,194,860	1,306	0.364
NAALADL2	3	174,156,363	1,367,065	1,198	0.456
CCSER1	4	91,048,686	1,474,378	1,398	0.441
PDE4D	5	58,264,865	1,553,082	1,166	0.458
GMDS	6	1,624,041	621,885	399	0.441
PARK2	6	161,768,452	1,380,351	1,296	0.555
IMMP2L	7	110,303,110	899,463	1,028	0.444
PTPRD	9	8,314,246	2,298,477	1,264	0.309

PRKG1	10	52,750,945	1,307,165	781	0.318
GPHN	14	66,974,125	674,395	291	0.306
WVVOX	16	78,133,310	1,113,254	1,319	0.541
MACROD2	20	13,976,015	2,057,827	3,039	0.605
DMD	X	31,115,794	2,241,764	789	0.328
DIAPH2	X	95,939,662	920,334	331	0.381

We also noted that 4 other significantly deleted genes (STS, HDHD1, LRRN3 and LINC00290), though not fulfilling the length criteria above have a particularly high proportion of deletion SVs between 20kb and 1 megabase (over 60%) and hence were also marked as fragile:

Gene	Chr	Start position	Length (bases)	Total Disruptive SV Count	% of SV that are DELs (>20kb & <1MB)
LINC00290	4	181,985,242	95,060	64	0.641
LRRN3	7	110,731,062	34,448	70	0.686
STS	X	7,137,497	135,354	168	0.649
HDHD1	X	6,966,961	99,270	126	0.659

Two of these genes (STS and HDHD1) fall in a previously identified CFS region (FRAXB) and a third, LRNN3, falls in another known CFS region (FRAX7). The final one, LINC00290 is a long non-coding RNA with an unknown status as cancer driver.

17. Driver catalog construction

We created a catalog of each and every driver in our cohort across all variant types on a per patient basis. This was done in a similar incremental manner to Sabarinathan et al¹⁹ (N. Lopez, personal communication) whereby we first calculated the number of drivers in a broad panel of known and significantly mutated genes across the full cohort, and then assigned the drivers for each gene to individual patients by ranking and prioritising each of the observed variants. Key points of difference in this study were both the prioritisation mechanism used and our choice to ascribe each mutation a probability of being a driver rather than a binary cutoff based on absolute ranking.

The four detailed steps to create the catalog are described below:

1. Create a panel of driver genes for point mutations using significantly mutated genes and known drivers

We created a gene panel using the union of

- Martincorena significantly mutated genes¹⁶ (filtered to significance of $q < 0.01$)
- HMF significantly mutated genes ($q < 0.01$) at global level or at cancer type level
- Cosmic Curated Genes¹² (v83)

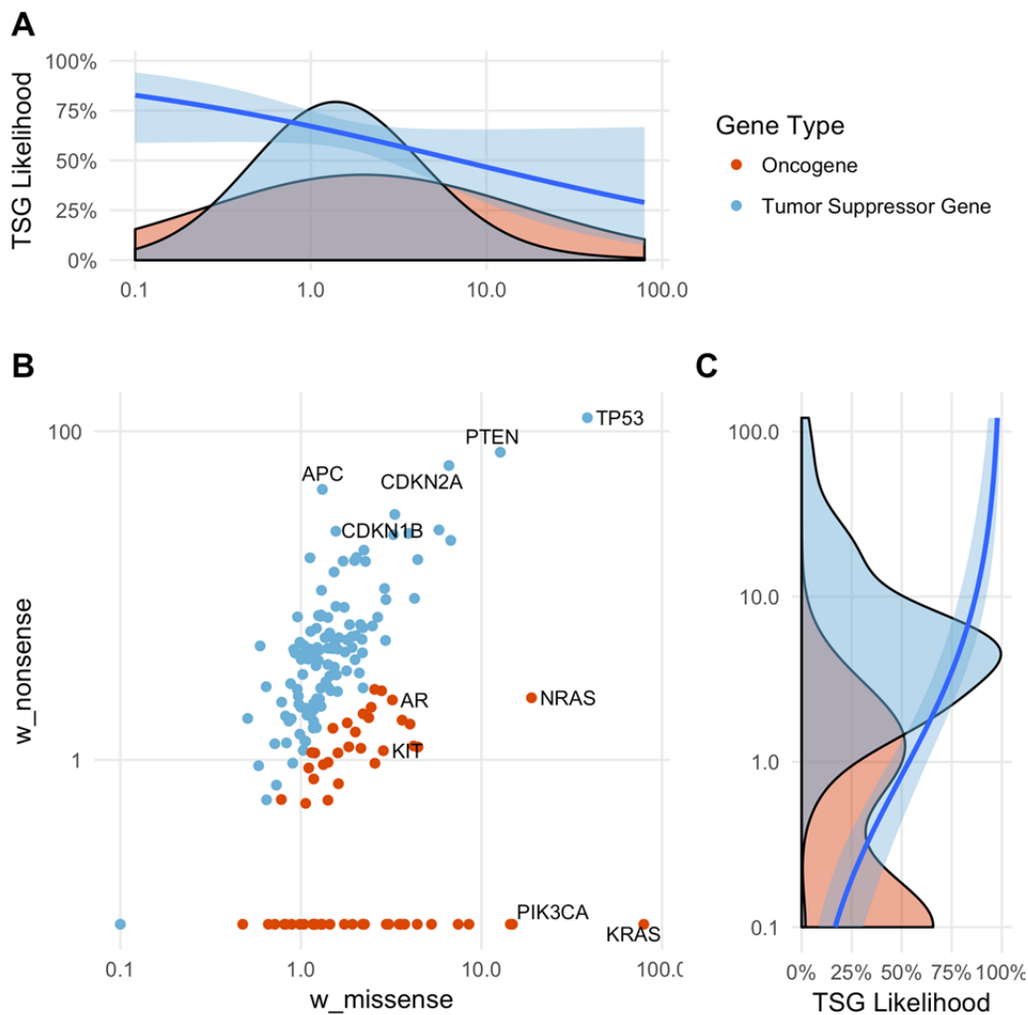
2. Determine TSG or Oncogene status of each significantly mutated gene

We used a logistic regression model to classify the genes in our pane as either tumor suppressor gene (TSG) or oncogene. We trained the model using unambiguous classifications from the Cosmic curated genes, i.e. a gene was considered either a Oncogene or TSG but not both. We determined that the dNdS missense and nonsense ratios (w_{missense} and w_{nonsense}) are both significant predictors of the classification. The coefficients are given in the table below.

	Estimate	Std. Error	z value	Pr(> z)
intercept	0.1830	0.3926	0.466	0.64106
w_{missense}	-0.6869	0.2643	-2.599	0.00936
w_{nonsense}	0.5237	0.1116	4.691	2.72e-06

We applied the model to all significantly mutated genes in Matincorena and HMF as well as any ambiguous Cosmic curated genes.

The following figure shows all genes that have classified using the logistic regression model. Figures A and C show the likelihood of a gene being classified as a TSG under a single variate logistic model of w_{missense} and w_{nonsense} respectively. Figure B shows the classification after the multivariate regression using both predictors.



3. Add drivers from all variant classes to the catalog

Variants were added to the driver catalog which met any of the following criteria

- All missense and inframe indels for panel oncogenes
- All non synonymous and essential splice point mutations for tumor suppressor genes
- All high level amplifications (min exonic copy number > 3 * sample ploidy) for both significantly amplified target genes and panel oncogenes
- All homozygous deletions for significantly deleted target genes and panel TSG (except for the Y chromosome as described before)
- All known or promiscuous inframe gene fusions as described above
- Recurrent TERT promoter mutations

4. Calculate a per sample driver likelihood for each gene in the catalog

A driver likelihood estimate between 0 and 1 was calculated for each variant in the gene panel to ensure that only excess mutations are used for determining the number of drivers in cancer cohort groups or at the individual sample level. High level amplifications, Deletions, Fusions, and TERT promoter mutations are all rare so were assumed to have a likelihood of 1 when found affecting a driver gene, but for coding mutations we need to account for the large number of passenger point mutations that are present throughout the genome and thus also in driver genes.

For coding mutations we also marked coding mutations that are highly likely to be drivers and/or highly unlikely to have occurred as passengers as driver likelihood of 1, specifically:

- Known hotspot variants
- Variants within 5 bases of a known pathogenic hotspot in oncogenes
- Inframe indels in oncogenes with repeat count < 8 repeats. Longer repeat count contexts are excluded as these are often mutated by chance in MSI samples
- Biallelic variants in tumor suppressor genes

For the remaining variants (non-hotspot missense variants in oncogenes and non-biallelic variants in TSG) these were only assigned a > 0 driver likelihood where there was a remaining excess of unallocated drivers based on the calculated dNdS rates in that gene across the cohort after applying the above rules. Any remaining point mutations were assigned a driver likelihood between 0 and 1 using a bayesian statistic to calculate a sample specific likelihood of each gene based on the type of variant observed (missense, nonsense, splice or INDEL) and taking into account the mutational load of the sample. The principle behind the method is that the likelihood of a passenger variant occurring in a particular sample should be approximately proportional to the tumor mutational burden and hence variants in samples with lower mutational burden are more likely to be drivers.

The sample specific likelihood of a residual excess variant being a driver is estimated for each gene using the following formula:

$$P(\text{Driver}|\text{Variant}) = P(\text{Driver}) / (P(\text{Driver}) + P(\text{Variant}|\text{Non-Driver}) * (1-P(\text{Driver})))$$

where $P(\text{Driver})$ in a given gene is assumed to be equal across all samples in the cohort, ie:

$$P(\text{Driver}) = (\text{residual unallocated drivers in gene}) / \# \text{ of samples in cohort}$$

And $P(\text{Variant}|\text{Non-Driver})$, the probability of observing n or more passenger variants of a particular variant type in a sample in a given gene, is assumed to vary according to tumor mutational burden, and is modelled as a poisson process:

$$P(\text{Variant}|\text{Non-Driver}) = 1 - \text{poisson}(\lambda = \text{TMB}(\text{Sample}) / \text{TMB}(\text{Cohort}) * (\# \text{ of passenger variants in cohort}), k=n-1)$$

All counts reported in the paper at a per cancer type or sample level refer to the sum of driver likelihoods for that cancer type or sample.

18. Driver co-occurrence analysis

To examine the co-occurrence of drivers, the driver-gene catalog was filtered to exclude fusions and any driver with a driver likelihood of < 0.5 . Separately for each cancer type, every pair of driver genes was tested to see whether they co-occur more or less frequently than expected if they were independent using Fisher's Exact Test. The results were adjusted to a FDR using the number of gene-pair comparison being tested in each cancer type cohort. Gene pairs with a positive correlation which were on the same chromosome were excluded from the analysis as they are frequently co-amplified or deleted by chance.

19. Actionability analysis

To determine clinical actionability of the variants observed in each sample, we mapped all variants to 3 external clinical annotation databases

- OncoKB⁷ (download = 01-mar-2018)
- CGI⁶ (update: 17-jan-2018)
- CIViC⁵ (download = 01-mar-2018)

In order to be able to aggregate and compare this data, we have mapped each of the databases to a common data model using the following rules:

1. Level of evidence mapping

The 3 databases we used in this study define different level for evidence items, depending on evidence strength. In order to be able to aggregate and compare this data, we have mapped the CGI and OncoKB evidence levels on the CIViC evidence levels defined at: <https://civicdb.org/help/evidence/evidence-levels>.

HMF	CIViC	CGI	OncoKB
A	A	FDA guidelines, NCCN guidelines, NCCN/CAP guidelines, CPIC guidelines, European Leukemia Net guideline	1 2 R1
B	B	Clinical trials, Late trials, Late trials,Pre-clinical	3 R2
C	C	Early trials, Case report	
D	D	Pre-clinical	4,R3

In this study we considered only A and B level variants. This classification roughly corresponds to the recently proposed ESMO Scale for Clinical Actionability of molecular Targets (ESCAT)²⁰ in the following way:

HMF A: ESCAT I-A+B (for on label) and I-C (for off-label)

HMF B: ESCAT II-A+B (for on label) and III-A (for off-label)

2. Response type Mapping

We also mapped response type to a common data model. First we filtered out evidence items from the annotation databases that do not lead to clinical actionability (for example prognostic biomarkers). The remaining evidence items were mapped as either responsive or resistant based on the following rules:

HMF	CIVIC	CGI	OncoKB
Responsive	Sensitivity	Responsive	1 2 3 4
Resistant	Resistant or Non-Response	Resistant	R1 R2 R3

3. Mutation/Event type mapping

Each evidence item was mapped to HMF data as one of 4 event types according to the following criteria

HMF Event type	Matching Criteria
Somatic Point Mutation	HGVS / genomic coordinates converted to chromosome, position, ref and alt and mapped to exact variants in our database
Somatic Range Event	Matched to missense / inframe variants in Oncogenes and any non-synonymous variant in TSG contained within a defined range, either exon level, transcript level or specific coordinates. Where a transcript was not specified, the canonical transcript was always used to map coordinates
Somatic CNA	'Deletion' mapped to homozygous deletions and 'Amplification' mapped to high level amplification (>3x sample ploidy)
Fusion	Exact matching to an inframe fusion in our database. For OncoKB 'loss-of-function' fusions were excluded

A small number of items from CIVIC level B evidence level were deemed either not specific enough or insufficiently supportive of actionability for this study and were filtered:

- Evidence items supporting TP53, KRAS & PTEN as actionable
- Evidence items supporting actionability with 'chemotherapy' (ie. chemotherapy in general rather than a specific treatment), 'asprin' or 'steroids'

Finally, a number of suspicious fusions from each of the database were curated by either changing the 5' and 3' partners or filtered out altogether based on referring to the original evidence sources, specifically:

HMF Curation	CIVIC	CGI	OncoKB
Filtered Fusions	BRAF - CUL1	RET - TPCN1	
5' and 3' partners exchanged		ABL1 - BCR PDGFRA - FIP1L1 PDGFB - COL1A1	ROS1 - CD74 EP300 - MLL EP300 - MOZ RET - CCDC6

Some of the more complex event types from the 3 databases have not been fully interpreted and have been excluded from this analysis.

4. Cancer type mapping

Each evidence event mapped was also determined to be either on-label (ie. evidence supports treatment in that specific cancer type) or off-label (evidence exists in another cancer type) for each specific sample. To do this, we have annotated both the patient cancer types and the database cancer types with relevant DOIDs, using the disease ontology database available at: <http://disease-ontology.org>.

Patient cancer types from the HMF database were annotated according to the following table:

HMF tumor type	DOID
Biliary	4607
Bone/Soft tissue	201;9253
Breast	1612
CNS	3620;3070
Colon/Rectum	9256;219
CUP	-
Esophagus	5041;4944
Head and neck	11934;8618
Kidney	263;8411
Liver	3571
Lung	1324
Mesothelioma	1790
NET	-
Other	-
Ovary	2394
Pancreas	1793
Prostate	10283
Skin	4159
Stomach	10534
Urinary tract	3996
Uterus	363

Database cancer types were mapped to a DOID by automatically querying the ontology on the disease names. Some CIViC evidence items are already annotated with a DOID in the database, this was used if present. We also manually annotated with DOIDs some of the database cancer types that failed the automatic query:

cancerType	DOID	Ontology term
All Tumors	162	cancer
Any cancer type	162	cancer
B cell lymphoma	707	B-cell lymphoma
Biliary tract	4607	biliary tract cancer
Bladder	11054	urinary bladder cancer
Cervix	4362	cervical cancer
CNS Cancer	3620	central nervous system cancer
Dedifferentiated Liposarcoma	3382	liposarcoma
Endometrium	1380	endometrial cancer
Esophagogastric Cancer	5041	esophageal cancer
Gastrointestinal stromal	9253	gastrointestinal stromal tumor
Giant cell astrocytoma	3069	astrocytoma

Hairy-Cell leukemia	285	hairy cell leukemia
Head and neck	11934	head and neck cancer
Head and neck squamous	5520	head and neck squamous cell carcinoma
Hepatic carcinoma	686	liver carcinoma
Hepatocellular Mixed Fibrolamellar Carcinoma	0080182	mixed fibrolamellar hepatocellular carcinoma
Inflammatory myofibroblastic	0050905	inflammatory myofibroblastic tumor
Lung	1324	lung cancer
Lung squamous cell	3907	lung squamous cell carcinoma
Melanoma	8923	Skin melanoma
Mesothelioma	1790	malignant mesothelioma
Neuroendocrine	169	neuroendocrine tumor
Non-small cell lung	3908	non-small cell lung carcinoma
Ovary	2394	ovarian cancer
Pancreas	1793	pancreatic cancer
Renal	263	kidney cancer
Salivary glands	8850	salivary gland cancer
Stomach	10534	stomach cancer
Thymic	3277	thymus cancer
Thyroid	1781	thyroid cancer
Well-Differentiated Liposarcoma	3382	liposarcoma

In case a matching DOID was found for the disease, we annotated the disease with a DOID set consisting of: the disease DOID, all the children DOIDs and all the parent disease DOIDs.

A treatment is defined as on-label if any of the DOIDs of the patient cancer is present in the DOID set of the disease.

5. MSI actionability

Samples classified as MSI in our driver catalog were also mapped as actionable at level A evidence based on clinical annotation in the OncoKb database

6. Aggregation of evidence

For each actionable mutation in each sample, we aggregated all the mapped evidence that was available supporting both on-label and off-label treatments at an A or B evidence level. Treatments that also had evidence supporting resistance based on other biomarkers in the sample at the same or higher level were excluded as non-actionable.

For each sample we reported the highest level of actionability, ranked first by evidence level and then by on-label vs off-label.

20. Data availability

All data described in this study is freely available for academic use from the Hartwig Medical Foundation through standardized procedures and request forms which can be found at <https://www.hartwigmedicalfoundation.nl>. Briefly, a data request can be initiated by filling out the standard form in which intended use of the requested data is motivated. First, an advice on scientific feasibility and validity is obtained from experts in the field which is used as input by an independent Data Access Board who also evaluates if the intended use of the data is compatible with the consent given by the patients and if there would be any applicable legal or ethical constraints. Upon formal approval by the Data Access Board, a standard license agreement which does not have any restrictions regarding Intellectual Property resulting from the data analysis needs to be signed by an official organisation representative

before access to the data is granted. Raw data files will be made available through a dedicated download portal with two-factor authentication. As of August 2018, more than 30 national and international requests have already been granted through this mechanism.

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doi:10.1093/annonc/mdy263