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Clonal heterogeneity of acute myeloid leukemia treated with the IDH2 inhibitor

Enasidenib

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35 Mutations in the gene encoding isocitrate dehydrogenase 2 (IDH2) occur in several 36 types of cancer, including acute myeloid leukemia (AML). In model systems, mutant 37 IDH2 causes hematopoietic differentiation arrest. Enasidenib, a selective small-38 molecule inhibitor of mutant IDH2, produces a clinical response in 40% of treated 39 relapsed/refractory AML patients by promoting leukemic cell differentiation. Here, we 40 studied the clonal basis of response and acquired resistance to enasidenib treatment. 41 Using sequential patient samples, we determined the clonal structure of 42 hematopoietic cell populations at different stages of differentiation. Pre-therapy IDH2 43 mutant clones showed variable differentiation arrest. Enasidenib treatment promoted hematopoietic differentiation from either terminal or ancestral mutant clones; less 44 frequently, treatment promoted differentiation of non-mutant cells. Analysis of paired 45 46 diagnosis/relapse samples did not identify second site mutations in IDH2 at relapse. 47 Instead, relapse arose by clonal evolution, or selection, of terminal or ancestral clones, highlighting multiple bypass pathways that could potentially be targeted to 48 49 restore differentiation arrest. Mapping clonal structure in cell populations at different 50 stages of differentiation during therapy illustrates how different clones respond and 51 evolve during relapse.

52 Differentiation arrest is a common feature of many cancer cells. Though intratumoral clonal 53 heterogeneity is well documented, we lack a detailed understanding of how the complement 54 of driver mutations within a clone, contribute to differentiation arrest. Furthermore, though it 55 is clear that genetic and functional, intratumoral heterogeneity helps determine clinical 56 outcome to cancer therapy^{1,2}, few studies have investigated the relationship between clonal 57 structure and therapy response, particularly for therapies targeting oncogenic epigenetic 58 processes. With an average of 13 somatic mutations per patient, arranged in an limited number of clones³, AML is simpler than most cancers from a genomic context, providing a 59 potential paradigm to answer these questions. 60

Somatic mutations in conserved arginine residues R140 and R172 in *ISOCITRATE* DEHYDROGENASE 2 (*IDH2*) occur in 15-25% of AML patients⁴⁻⁶. The mutant proteins have neomorphic activity producing R-2-hydroxyglutarate (2-HG) that competitively inhibits αketoglutarate-dependent enzymes including the TET family of 5-methylcytosine (5mC) hydroxylases and Jumonji-C domain histone demethylases^{7,8}. This leads to DNA hypermethylation⁹, increased repressive histone methylation⁸ and impaired hematopoietic differentiation that is reversed by mutant IDH inhibition in model systems ^{8,10-12}.

We recently showed that enasidenib (AG-221/CC-90007), a first-in-class, allosteric inhibitor 68 of mutant IDH2 (mIDH2)¹³, reduces serum 2-HG, reverses DNA hypermethylation and 69 promotes hematopoietic differentiation in preclinical models^{13,14}. In a phase 1/2 clinical trial, 70 enasidenib monotherapy produced a response rate of 40.3% in relapsed/refractory AML 71 patients¹⁵. In most responding patients terminally mature blood cells were *IDH2* mutant. 72 consistent with response due to enasidenib-induced differentiation of *IDH*2 mutant cells¹⁶. 73 However, in 9/71 (12.6%) responding patients, IDH2 mutant cells were eliminated from 74 75 peripheral blood cells¹⁶. Failure to respond to enasidenib was associated with a higher comutational burden and NRAS mutations¹⁶. Finally, most patients who initially responded 76 77 eventually relapsed. These initial studies did not assess which clone(s) differentiated in 78 response to enasidenib and the clonal mechanism of acquired enasidenib resistance.

79 We have addressed these two questions by studying sequential samples from a subset of 80 trial patients. We showed marked variation in the degree of differentiation arrest of mutant 81 IDH2 clones. Restoration of differentiation by inhibition of mutant IDH2 was also clone 82 dependent, varying between patients, arising from either ancestral or terminal clones. In a 83 minority of patients differentiation occurred from wild type progenitor cells, consistent with 84 molecular remission in a subset of patient. Acquired resistance to enasidenib leading to 85 differentiation arrest and relapse did not occur by second site mutations in *IDH2*. Instead, 86 differentiation arrest was restored by multiple mechanisms through clonal evolution or clonal 87 selection.

88 Results

89 Patient cohort studied

The trial enrolled 176 relapsed/refractory IDH2 mutant AML patients¹⁵. Here, we studied a 90 91 cytogenetically and genetically representative subset of 37 patients enriched for enasidenib responders (30/37 responders, Supplementary Fig. 1a-c). An extended mutational profile 92 93 was determined in 33/37 patients by either whole exome sequencing (WES) (16/36 patients 94 at read depth of 19-843x, average 121x at loci where variants were called), or targeted 95 resequencing (17/36 patients) (Supplementary Tables 1 and 2). Compared to the entire 96 trial cohort, the patient cohort studied here had similar serum baseline 2-HG levels and the 97 mean level of 2-HG suppression was similar in the two cohorts (i.e. on-target response to enasidenib; Supplementary Fig. 1d-e). 98

99 Enasidenib rebalanced progenitor and precursor compartment sizes and restored 100 progenitor function

In AML there are two orthogonal potential hierarchies (**Supplementary Fig. 2a**). There is clonal hierarchy with an initiating clone, transitional and terminal clones. We refer to all nonterminal clones as ancestral clones. These mutant clones exist in a second hierarchy, a hematopoietic cell hierarchy. AML initiating mutations occur in stem, or long-lived progenitor, cells but initiating clones are not usually arrested in differentiation¹⁷⁻¹⁹. However, with

106 acquisition of additional, transforming mutations and epigenetic alterations, clones fail to 107 complete maturation. In the fully transformed state, haemopolesis in human AML is 108 dominated by expansion either of progenitor-like cells, presumably because of a differentiation block between progenitor and downstream precursor cells²⁰, or less 109 110 commonly, by precursor-like cells, presumably because of a differentiation block between precursor and mature cells²¹. In both cases, expanded leukemic progenitor-like, or 111 precursor-like, populations have functional leukemic stem cell activity^{20,21}. Thus, we set out 112 113 to address three questions: (i) where are individual clones arrested in the hematopoietic 114 hierarchy; (ii) which clone(s) responded to mutant IDH2 inhibition by differentiating; (iii) which 115 clone(s) were responsible for loss of response to enasidenib, after an initial response, and 116 by what mechanism?

117 The experimental approach is set out in Supplementary Fig. 2b. We first performed flow 118 cytometric quantitation of hematopoietic stem/progenitor (Lin⁻CD34⁺CD117⁺), precursor (Lin⁻ CD34⁻CD117⁺) and mature myeloid cell (Lin⁻CD34⁻CD117⁻) populations²¹ at trial entry in 15 119 120 patients (Fig. 1a-b and Supplementary Fig. 3a-b). Sizes of individual stem/progenitor 121 populations within Lin⁻CD34⁺ cells were also quantitated. 11/15 patients had abnormally 122 expanded progenitor-like compartments (mainly LMPP-like and GMP-like; termed progenitor 123 AML) and 4/15 abnormally large myeloid precursor-like populations (termed precursor AML). The ratio of progenitor: precursor AML is consistent with previous studies^{20,21}. 124

125 Next, we analysed the bone marrow (BM) stem/progenitor/precursor populations in 5 126 patients who achieved complete remission (CR) with enasidenib (Fig. 1b-c). In all 5 there 127 was near normalization of the sizes of the stem/progenitor compartments. Pre-treatment, 2 128 patients had pathologically expanded LMPP- and GMP-like progenitor populations (#201-129 023, #201-011) and 2 patients had expanded myeloid precursor compartments (#201-010 130 and #203-002). Functionally, Lin CD34⁺ progenitor cells from these 5 patients in CR formed 131 myeloid/ erythroid colonies nearly as efficiently as normal cells, in contrast to cells from 132 patients who did not achieve CR (Fig. 1d)(Supplementary Fig. 3c-d). Thus, enasidenib

- 133 therapy rebalanced the sizes of hematopoietic stem/progenitor/precursor/mature populations
- at CR with reacquisition of normal myeloid progenitor function.

135 Wild type hemopoiesis occurs occasionally with enasidenib therapy

136 Next, we investigated the clonal basis of differentiation. In principle, enasidenib could have, 137 directly or indirectly, restored differentiation from either wild type cells, ancestral clones or 138 terminal clones in a clonal hierarchy (Fig. 2a). We established the clonal basis of response 139 in 6 patients using samples taken at multiple time points before and through treatment, 140 including relapse (Fig. 2b). We used WES and karyotype of bone marrow mononuclear cells 141 (BMMNCs) to determine chromosomal copy number and mutational changes 142 (Supplementary Tables 2-3). Next, we used WES data to design patient-specific mutation 143 panels to test variant allele frequencies (VAFs) of mutations in unsorted BMMNCs and flow-144 cytometric sorted hematopoietic stem/progenitor/precursor/mature cell populations and 145 genotype flow-cytometric sorted single cells and hematopoietic colonies derived from single 146 cells (Supplementary Table 3). A combination of all these data was used to establish clonal 147 structures. Details on setting false positive and negative thresholds in single cell genotyping 148 (SCG) are presented in Methods.

149 In our initial study, 9 out of 29 patients who achieved CR, for whom samples were available, 150 had loss of mutant IDH2 peripheral blood cells (complete molecular remission in peripheral blood¹⁶. However, it was unclear if these differentiated blood cells were truly wild type, or 151 152 from a genetically mutant clone(s) that just lacked mutant *IDH2*? We studied this question in 153 a patient with mutations in IDH2 (I), PEX26 (P), FEZ2 S208T (F), DNMT3B (D), ZCCHC1 154 (Z), NPM1 (N), and ELMO3 (E) in AML blasts pre-enasidenib (#201-022, Fig. 2c) who 155 achieved mutant IDH2 molecular remission. Pre-therapy, imputation from variant allele 156 frequency (VAF) suggested presence of wildtype cells and three possible mutant clones: a 157 clone with *IDH*² mutation alone (*I*), a clone with genotype *IPFDZN* and a minor clone that 158 was either *IPFDZNE* or *IE* (Supplementary Fig. 4a-c), though the exact clonal structure could not established unambiguously (Supplementary Fig. 4d-h). At CR, the VAF of all 159

mutations was <1.6% (**Fig. 2c, Supplementary Table 4 for depth of sequencing**). Concordantly, at CR the majority (94.6%) of 111 individually genotyped hematopoietic colonies did not contain any mutations present pre-therapy (**Fig. 2d**). Functionally, the colonies produced a normal ratio of myeloid to erythroid colonies consistent with a wild-type genotype (**Fig. 2e**). Thus, in a minority of patients, enasidenib therapy resulted in restoration of wild type terminal blood cell production and progenitor function from wild-type cells.

166 Enasidenib restored differentiation from ancestral and terminal clones in a clone 167 dependent manner

168 In one patient enasidenib promoted differentiation from an ancestral clone (#201-023, Fig. 169 3). WES and targeted resequencing of BMMNCs pre-enasidenib detected mutations in 170 SRSF2 (S), IDH2 (I), ASXL1 (A), GATA2 (G) and two mutations in RUNX1 (R, r) 171 (Supplementary Tables 2-3). Pre-therapy imputation of clonal structure based on VAF in 172 unsorted BMMNCs by exome and targeted resequencing suggested initial acquisition of 173 SRSF2 mutation (clone S) followed by an IDH2 mutation (clone SI) followed by acquisition of 174 ASXL1 (A), the two RUNX1 and GATA2 mutations (Supplementary Fig. 5a and 175 Supplementary Tables 2-3). At CR, only mutations SRSF2 (S), IDH2 (I), ASXL1 (A) were 176 detected in mature cells supporting the existence of an SIA clone that preferentially 177 completes terminal maturation in presence of enasidenib (Supplementary Fig. 5a-c).

178 To clarify clonal structure, and position clones within the haemopoietic hierarchy, we 179 performed targeted resequencing for driver variants in 63 single cells and cell populations, 180 from flow-cytometric sorted progenitor and mature myeloid cell compartments, from both 181 pre-therapy and CR samples (Fig. 3a-b and Supplementary Table 3). This confirmed a 182 linear clonal evolution pattern: clone SI preceded clone SIA, followed by clones that 183 sequential acquired RUNX1 mutations (clones SIAR and SIARr) and finally the terminal 184 clone acquired a GATA2 mutation (SIARrG). The mutational profile in 4 single cells did not fit 185 into this clonal evolution pathway. We detected a single cell with an ASXL1 mutation (clone 186 A) and three single cells with both ASXL1 and SRSF2 mutations (clone AS). In three cells

with genotype AS, we detected allele dropout (ADO) of the *IDH2* allele in two out of three cells. Thus, we are unable to determine the mutational state of *IDH2* in those two cells. In contrast, in twelve cells of the SI clone, ADO of the *ASXL1* allele was detected in only three out of twelve cells (**Supplemental Fig. 12a**). Thus, though our results do not exclude a rarer, parallel clonal evolution pathway where clones A and AS exist and failed to acquire mutations in the order shown in the main pathway, they also are consistent with these cells being part of the main clonal evolution pathway.

Pre-therapy, 90% of BMMNCs were progenitors (LMPP and GMP) (Fig. 1c and 3d) and 90-194 195 100% of these leukemic progenitors were the SIARrG clone (Fig 3c). Thus, the SIARrG 196 clone is arrested in differentiation at the progenitor stage and expands to dominate the 197 marrow. Less than 10% of BMMNCs were mature myeloid cells (Fig 1c and Fig 3d). We 198 were only able to genotype 8 mature myeloid cells pre-therapy (Fig. 3c) and they are 199 composed of a mixture of wild type cells and of cells with genotypes A, SI, AS and SIA. At 200 CR, the mature myeloid cell compartment comprised 60% of BMMNCs (Fig 1c and Fig 3d) 201 and 85% of mature cells have the SIA genotype (Fig. 3d and Supplementary Fig. 5b, 202 Supplementary Table 3).

The progenitor compartment was only 20% of BMMNCs at CR (**Fig. 1c and Fig 3d**) and composed of mixed ancestral clones SI, SIA, SIAR but not the terminal clone SIARrG. Concordantly, the majority of colonies generated by progenitors at CR were "SIA", with a minority of the SI and SIA genotype (**Fig. 3e**). The ratio of myeloid:erythroid colonies was within normal limits (**Fig. 3f**) Taken together, this data demonstrated a complex, clonedependent pattern of enasidenib-induced differentiation with mature myeloid cell production sustained principally by a self-renewing ancestral *SIA* clone.

In four patients, differentiation of mature cells was principally seen from terminal clones (Fig.
2b). In patient #201-011, WES (Supplementary Table 2) and targeted resequencing of
BMMNC revealed two mutations in *DNMT3A* and mutations in *IDH2*, *ASXL1* and *XPO1* pretherapy that persisted at different VAFs at CR (Supplementary Table 3, Supplementary

214 Fig. 6a). However, it was not possible to impute the clonal structure from the VAF 215 (Supplementary Fig. 6b). We genotyped 110 single cells, pre-therapy and at CR (Fig. 4a). 216 This revealed an initiating DNMT3A clone (clone D) that acquired an IDH2 mutation (clone 217 DI) (Fig. 4b). Subsequently, there is a branching clonal structure with two terminal clones; 218 one acquired an XPO1 mutation (clone DIX), whereas the other acquired two mutations, a 219 second DNMT3A mutation and an ASXL1 mutation (clone DIdA). SCG suggested the DIdA 220 clone may have arisen by convergent evolution through intermediate DId and DIA clones. 221 There is a caveat with this interpretation as ADO was detected in 6 out of 7 DIA clone cells 222 (Supplemental Fig. 12d). In the DId cells, though there were no heterozygous germline 223 single nucleotide polymorphisms in the ASXL1 gene, the estimated ADO frequency of the 224 ASXL1 allele was 12.1% (Supplemental Fig. 12e). Thus, it is also possible that the DIdA 225 clone may also have arisen through just one mutational pathway.

Pre-enasidenib, 89% of BMMNCs were leukemic progenitors (LMPP and GMP) virtually exclusively composed of the DIX clone (**Fig. 4c-d**). A small mature myeloid population is present pre-enasidenib composed of the DIdA clone (**Supplementary Fig. 6c**). These observations suggest that the *IDH2* mutation in the context of the DIdA clone is not fully effective at imposing a complete differentiation block, whereas the same *IDH2* mutation in the context of the DIX clone fully arrests at a progenitor stage.

232 At CR, 82% of BMMNCs were composed of mature myeloid cells, 85% of which were a mix 233 of two terminal branching clones DIdA (54%) and DIX (31%) suggesting enasidenib 234 promoted differentiation from both terminal clones (Fig. 4c-d, Supplementary Table 3). To 235 address which progenitors contribute to mature cell output we genotyped single flow 236 cytometric sorted progenitors. The DIdA clone dominated mature GMP, CMP and MEP 237 progenitor compartments (Fig. 4c-d). In contrast, clone DIX was detected only in the more 238 immature LMPP progenitor compartment. GMP, CMP and MEP are more clonogenic than the LMPP^{20,22} and concordantly, ~95% of colonies had the DIdA genotype (Fig. 4e) that 239 240 were myeloid-biased (Fig. 4f). Interestingly, there is a substantial decrease in the size of the

LMPP compartment at CR compared to pre-therapy (Fig. 4d) (1240-fold decrease within the
 Lin⁻ compartment and 81-fold within the CD34⁺ compartment).

In three additional patients the terminal clone contributed to mature myeloid cells at CR based on imputed clonal structures, genetic analysis of mature myeloid cells at CR (patient #201-010, **Supplementary Fig. 6d-e**) and genotyping of myeloid colonies at CR (patient #201-027, **Supplementary Fig. 7a-d**; patient #201-006 **Supplementary Fig. 7e-h**). Depth of coverage for each of the mutations in all three patients is in **Supplementary Table 4**.

In summary, enasidenib therapy provides relief of differentiation arrest at a progenitor-like or precursor-like stage, normalizing the sizes of these abnormally expanded compartments. The ability of mutant *IDH2* to impose differentiation block is dependent on the context of coassociated mutations within a clone. Consequently, efficacy of enasidenib-induced differentiation is also is likely to dependent mutational landscape within a clone.

Relapse of IDH2 mutant patients on enasidenib occurs by clonal selection/evolution and not second site mutations in *IDH2* gene

255 Although responding patients have a median survival of 18-21 months, many patients relapse¹⁵. To study mechanisms leading to relapse we measured 2-HG levels in 16 patients 256 257 at diagnosis and relapse, mutational profiles in 12 patients (by WES in 11 cases and 258 targeted sequencing in the other patient) and performed karyotype analysis in 11 subjects 259 (Fig. 5a-b and Supplementary Table 5). We did not detect second site IDH2 mutations at 260 relapse in any patient but instead documented 7 patterns of clonal evolution/selection with 261 acquisition of recurrent AML-associated genetic changes (Fig. 5a). These are recurrent 262 missense mutations in myeloid malignancy, or nonsense and frameshift mutations in 263 cancers, as documented in the COSMIC database (http://cancer.sanger.ac.uk/cosmic). For 264 patient #201-007, mutations were detected prior to relapse, but increased in frequency at 265 relapse (Supplementary Table 3).

266 In 14/16 patients 2-HG levels remained suppressed between best response (CR or PR) and 267 relapse suggesting drug was on target in suppressing neomorphic enzyme function (Fig. 268 5b). However, in 2 patients (#201-014 and #201-022) rising 2-HG levels and BM leukemic 269 cells (blasts) were seen (Fig. 5c-d). Exome sequencing revealed IDH1 R132C/H mutations 270 albeit accompanied by other genetic abnormalities, some of which are recurrent in AML 271 (point mutations in RUNX1, NPM1 and t(3:12)) (Fig. 5e-f). These IDH1 mutations were 272 previously undetectable by high depth NGS (10000x) pre-enasidenib therapy. Surprisingly, 273 in both cases, the VAF indicates that *IDH1* mutations were present in *IDH2* mutant clones.

274 Relapse also associated with increasing VAF of oncogenic gain of function mutations in 275 genes encoding cytokine receptors CSF3R (patient #104-021, Supplementary Fig. 8a-b) 276 and FLT3 (patients #201-013, #201-004 Supplementary Fig. 8c-d and #201-007 277 Supplementary Fig. 9f-h) and predicted loss of function mutation in the negative regulator 278 of cytokine signalling CBL (#201-004 Supplementary Fig. 8d). For patient #104-021 we 279 could not resolve the clonal structure from targeted resequencing of BMMNC 280 (Supplementary Fig. 8b, Supplementary Table 2) but genotyping of 214 single cells pre-281 enasidenib and at relapse (Fig. 6a) demonstrated an initiating DNMT3A/ IDH2 mutant clone 282 that spawned the major clone with a recurrent U2AF1 mutation (DIU clone). A minor DIUF 283 clone with a D200E variant in the FLT3 (not previously described in AML) was also present 284 pre-enasidenib. At relapse, the major clone in the expanded arrested LMPP and GMP 285 compartments had acquired an oncogenic T618I mutation in the cytokine receptor CSF3R 286 that is well described in myeloid leukemias²³ (*DIUC clone*). DIUC further evolved, acquiring a 287 variant in NFKB1 that has not been described previously in AML. Mutations in both CSF3R 288 and *NFKB1* were detectable at threshold of sensitivity pre-enasidenib (**Supplementary Fig.** 289 8a).

290 Relapse, and re-imposition of differentiation block, was also associated with previously 291 described mutations in hematopoietic transcription factors in myeloid cancers. These 292 included frame shift mutations in *RUNX1* (**Fig. 5e and Supplementary Fig. 8e** one of which

has been previously described in AML (*RUNX1* F416fs)²⁴) and *BCORL1*²⁵; non-synonymous
variants, in the DNA- and protein-partner binding N-terminal zinc finger of *GATA2*(Supplementary Fig. 8d)^{26,27} and in one of the zinc fingers of *BCL11A*²⁸ (Supplementary
Fig. 8c). In all patients these mutations were not detected pre-enasidenib.

297 Deletion of all (monosomy 7), or part (del 7q) of chromosome 7 is common in myeloid malignancy²⁹. Chromosomal abnormalities were present in 18% of the enasidenib cohort¹⁵ 298 299 and 20% in the cohort studied here (Supplementary Fig 1). Del 7 was detected in 4 out of 300 12 patients who relapsed (#201-010, #201-007, #201-019 and #201-003) (Fig. 5a, 301 Supplementary Table 5) but are not enriched in relapsed patients (Amatangelo, M., 302 Thakurta, A., unpublished data). In all four patients it was detected either cytogenetically, or 303 by WES, pre-enasidenib therapy (Supplementary Tables 2 and 5). In three out of the four 304 cases the del 7q clone was selected at relapse (Supplementary Fig. 9a-e, k and I). In one 305 case where it was not, it was the dominant clone pre-therapy and at relapse 306 (Supplementary Fig. 9g-j).

307 Clonal evolution at relapse also highlighted variants in genes less well studied in AML 308 including NFKB1 M216I (Fig. 6a-d), DDX1 G699A (Supplementary Fig. 8c), MTUS1 309 Q781H (Supplementary Fig. 9f-i), DHX15 R222G and DEAF1 N372K (Fig. 6e-h). Of these, 310 acquisition of latter two variants by patient #201-011 at relapse is worthy of comment. Pre-311 enasidenib the patient had expanded LMPP and GMP populations composed of an arrested 312 DIX clone (Fig. 4b). At CR the DIX clone was able to differentiate but only persisted within 313 the LMPP compartment (Fig. 4d). At relapse, we detected 30 cells with DIX mutations that 314 acquired a missense mutation in the DExD/H-box helicase DHX15 R222G (mutation H), 7 315 cells with the DIX mutations that had acquired a variant in the transcription factor DEAF1 316 N372L (variant F) and 12 cells with the DIXHF variants (Fig. 6e-f). However in 6 out of 7 317 DIXF cells there was ADO for the DHX15 allele with the DHX15 R222G mutation and 318 therefore we are unable to determine whether the DHX15 R222G is present in those cells 319 (Supplementary Fig. 12c).

Acquisition of the additional *DHX15* and *DEAF1* mutations was associated with differentiation arrest, and a re-expansion of LMPP and GMP progenitor compartments, comprising 89% of MNCs. 83% of LMPP cells were composed of the DIXH and DIXHF clones whereas in the GMP-like compartment there was a more even contribution by the *DIXHF, DIXH* and *DIXF* clones (**Fig. 6g-h**).

325 Finally, both DHX15 and DDX1 regulate RNA splicing. Human DHX15 is structurally closely related to its yeast homologue, Prp43^{30,31}. Both proteins have been shown to contribute to 326 disassembly of splicesomes, efficient debranching and turnover of excised introns^{32,33}. 327 DHX15 R222G mutations have been previously described in AML^{34,35}. To determine if 328 acquisition of DHX15 R222G results in altered splicing we performed RNA-Seg of AML 329 blasts at relapse from patient #201-011 and compared splicing to AML cells from the same 330 331 patient that were wild type for DHX15 at trial entry (Supplementary Fig. 8f). In cells 332 expressing DHX15 R222G, there were alterations in exon skipping and intron retention 333 compared to cells wild type for DHX15. DDX1 is a DEAD-box RNA helicase with 5' single 334 stranded RNA exonuclease activity postulated to have multiple roles in RNA metabolism³⁶. 335 DDX1 G699A has not been previously described as a cancer-associated mutation nor has 336 its impact on RNA splicing been studied. Our data shows an increase in intron retention and 337 use of alternative 5' and 3' splice sites, and a decrease in spliced exons in mutant compared 338 to wild type AML cells from the same patient (Supplementary Fig. 8g).

339 Discussion

This study of clonal response and acquired resistance in sequential paired samples from AML patients treated with an IDH2 inhibitor extends prior preclinical studies^{13,14}. Preenasidenib there were complex patient- and clone-specific patterns of differentiation arrest. At CR, wildtype dominated cellular reconstitution was less common but does occur. More commonly, enasidenib causes a clone-specific differentiation response, either from ancestral or terminal clones, leading to near normalization of the sizes and functionality of progenitor and precursor hematopoietic compartments with altered clonal mix. Acquired resistance was

never due to a second site mutation in the same *IDH2* allele but instead due to either clonal
evolution or clonal selection. At least 7 different mutational mechanisms led to re-imposition
of differentiation arrest (Supplementary Fig.10).

350 In most patients enasidenib was unable to promote terminal differentiation and eradication of 351 IDH2 mutant clones; ancestral and/or terminal clones remained at CR. In patients with 352 restitution of wild type hemopoiesis, we infer that enasidenib most likely promoted terminal 353 differentiation of arrested self-renewing IDH2 mutant cells, allowing normal cells to dominate 354 hemopoiesis. Longer term clinical follow up of molecular CR patients will determine if 355 molecular CR patients have a better clinical outcome and if they relapse, the clonal origin of relapse. More generally, understanding the molecular mechanisms of relief from 356 357 differentiation arrest by IDH2 inhibitors will require in depth study of changing patterns of 358 epigenetic marks and transcriptional programmes within highly purified, clone-specific, 359 hematopoletic stem, progenitor and precursor populations, before and after drug exposure, 360 as transcriptional and epigenetic profiles are so highly plastic through differentiation.

Drug resistance to targeted cancer therapy arises by multiple mechanisms. Resistance to kinase inhibitors in AML^{37,38}, chronic myeloid leukemia³⁹, chronic lymphoid leukemia⁴⁰ and lung cancer^{41,42} often involves second site mutations in the mutant allele modulating drug or substrate binding or copy number changes of the mutant kinase. We did not observe this in enasidenib treated patients.

366 Acquired resistance led to IDH2 could arise by either epigenetic or genetic mechanisms or a 367 combination of the two. In most patients 2HG remained suppressed at relapse suggesting 368 that enasidenib remained on target and relapsed clones were not dependent on mutant 369 IDH2. In hematologic malignancies, genome-wide epigenetic variation (DNA methylation for 370 example) can be several orders more variable than genetic change⁴³⁻⁴⁵, is somatically 371 heritable, and subject to selection. Locus-specific DNA methylation (epiallele) variation 372 shows dynamic change in AML between diagnosis and relapse and can occur with distinct 373 kinetics, such that some patients have a high epiallele diversity and low somatic mutation

burden and vice versa⁴⁵. In melanoma, resistance to a B-RAF inhibitor arose in rare cells
through stochastic, transient variation in gene expression that was selected for by therapy⁴⁶.
This is consistent with prior work on chromatin mediated drug resistance in cancer cell
lines⁴⁷.

378 Clones acquiring gene mutations, or grosser genetic changes, have previously been reported in therapy resistant chronic lymphocytic leukemia⁴⁸ and medulloblastoma⁴⁹. 379 380 Acquisition of an *IDH1* mutation in two patients is an example of how this may occur. Here, 381 these AML propagation is likely to be highly dependent to high 2HG. 2HG addiction may be 382 AML cell autonomous or alternatively, the 2HG requirement may be in BM niche supporting 383 cells or other non-AML cell populations. AML clone-specific and non-AML cell specific 384 analysis of the impact of 2HG on epigenetic and transcriptional programs and metabolism of cells⁵⁰ is needed to understand this dependency. Other examples genetic changes leading 385 386 to enasidenib resistance include gain of function mutations in proliferative cytokine signalling 387 pathways and loss of or altered function in transcriptional regulators of hemopoiesis. However, mutations in the RAS pathway that are correlate with failure of initial response¹⁶ 388 389 were not associated with acquired resistance.

390 We also detected variants at relapse not previously well studied in AML. An example of this is the DHX15 R222G mutation, recently described in RUNX1-RUNXT1 AML³⁴. The yeast 391 392 homologue of DHX15, Prp43, regulates RNA splicing and ribosome biogenesis. Loss of wild type DHX15 and overexpression of mutant DHX15 increases alternative splicing. In contrast, 393 394 a role for DEAF1 has not been previously published in normal or malignant hemopoiesis. 395 Curiously DEAF1 is a paralog of the transcription factor, RUNXT1. DEAF1 is expressed throughout hemopoiesis but particularly in GMP and AML blasts²¹. In non-hemopoetic 396 397 tissues it binds to LMO4, a member of the LMO transcriptional adapter protein family. In 398 blood cells, LMO2, a closely related LMO family member, partners transcription factors 399 including GATA, E-box proteins and LDB1 to form regulatory and oncogenic protein 400 complexes. Thus, the role of DEAF1 in hemopoiesis and its interaction with DHX15 merits

401 further study. More broadly, a deeper mechanistic understanding of how wild-type IDH2 402 promote hematopoietic differentiation, which is currently poorly understood, will also 403 increase our understanding of how bypass pathways could re-impose differentiation block, 404 for example by altering transcriptional programmes, as seen in BET inhibitor therapy 405 resistance⁵¹.

406 Finally, this study demonstrates how any cancer therapy alters clonal structure across a fully 407 transformed and pre-malignant cellular hierarchy. By defining clonal structures and mapping 408 where clones were arrested across differentiation, in purified hematopoietic compartments, 409 we obtained a previously unavailable view of where different clones were arrested. This 410 provides the necessary information to study why clones are arrested at different stages of 411 differentiation. Furthermore, analysis of sequential samples through therapy shows how 412 clones differentially responded to therapy. This provides the basis to study clone specific 413 relief of IDH2 inhibition. More generally, our approach could be applied to any cancer 414 therapy, where single cell suspensions and purification of cells at different stages of 415 differentiation is possible. This would then provide a clone specific understanding of how 416 therapy alters clonal structure through a cellular hierarchy. Our approach also provides 417 insights towards a rational basis for combination therapies to reduce drug resistance.

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438 Author contributions:

L.Q., M.D.D. designed/ performed experiments, analysed data; A.K., M.M., M.A., B.S., C.Q., M.H., C.W., V.S., S.AI. performed experiments/ analysed data; M.S.V. and G.S.V. analysed data; M.A., A.S., A.P., K.Y., S.Ag., S.dB., R.L.L., E.S., K.M., A.T. provided reagents/samples/clinical data; O.A.B., S.dB., A.T., R.L.L., V.P.-L. and P.V. designed the experiments/ analysed the data. L.Q. and P.V. wrote the manuscript. All authors edited the manuscript.

445

446 **Competing interests**:

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Data and materials availability: Exome sequencing data has been deposited in
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 ArrayExpress under accession E-MTAB-6660.

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- 572

573 Figure Legends

574 Figure 1. Enasidenib treatment induces differentiation of AML progenitor and 575 precursor cell populations and restores progenitor function.

576 a) Top, immunophenotyping of hematopoietic stem/progenitor/precursor and mature cell 577 populations in AML bone marrow (BM) samples pre-treatment showing expanded progenitor 578 (n=11 biologically independent samples) or precursor (n=4 biologically independent 579 samples) populations with normal BM (n=8 biologically independent samples). Below, 580 detailed composition of stem/progenitor compartments in AML pre-treatment (n=15 581 biologically independent samples) and normal BM (n=12 biologically independent samples). 582 Error bars in normal BM= 95% confidence interval. HSC: hematopoietic stem cell, MPP: 583 multipotent progenitor, LMPP: lymphoid-primed multipotent progenitor, CMP: common 584 myeloid progenitor, MEP; megakaryocyte-erythroid progenitor; GMP: granulocyte-585 macrophage progenitor.

b) Top, schematic representation of flow cytometric approach and sequential gates used to analyse samples in (a). Lin⁻, lineage negative; BMMC, bone marrow mononuclear cells. Bottom, example of flow plots from a representative sample prior to enasidenib treatment (Pre-ENA) and at complete remission (CR) in patients with expanded progenitor-like populations (#201-023; left, experiment performed once) or expanded myeloid precursor-like population (#201-010; right, experiment performed once). Numbers shown within the gate indicate percentage of the corresponding cell population compared to all cells in the plot.

c) Top, immunophenotyping of hematopoietic cell populations in normal BM (left, as in (a))
and in samples from 5 patients (#201-023, #201-011, #201-022, #201-010, #203-002) preENA, at intermediate time points during treatment, and at CR. Bottom, sizes of stem and
progenitor compartments. Abbreviations and error bars in normal BM are as in (a). C=cycle,
D=day.

d) Number of mixed erythroid-myeloid colonies (GEM), granulocyte-macrophage (GM), granulocyte (G), macrophage (M) and erythroid (E) colonies produced per 100 plated flowsorted CD34⁺ cells from normal BM (n=4 biologically independent samples), enasidenibtreated patients in CR (n=5 biologically independent samples) and enasidenib-treated patients not in CR (n=3 biologically independent samples). Patient samples were plated with addition of Enasidenib (1 μ M) to semi-solid media. Error bars= standard error of the mean. Pvalues determined by 2-sided Student's paired t-test.

Figure 2. Differentiation response arising from wild-type cells in patients treated with enasidenib.

a) Schematic representation of varying possible clonal responses to enasidenib. Four
mutations (A, B, C and D) are present in four clones that are arranged in a branching
structure. A differentiation response to enasidenib treatment could potentially occur from
either wild type cells or from ancestral or terminal clones.

b) Summary of the type of differentiation response (from either wild type cells, ancestral or
terminal clones) in samples from 6 patients.

c) Variant allele frequencies (VAF) of the indicated mutations in AML blasts of patient #201022 prior to enasidenib treatment (Pre-ENA) and in peripheral blood mononuclear cells
(PMNC) at CR, as assessed by targeted re-sequencing.

d) Clonal contribution to colony output from the CR sample from patient #201-022, as a
percentage of all individually picked colonies genotyped. Clones were identified as wild type
(WT), carrying the *FEZ2* P118S mutation (F), or carrying the *FEZ2* P118S and *DNMT3B*

N738S mutations (FD). Lineage affiliations of the colonies are as in Fig. 1d. Numbers next
to the bars indicate the number of colonies analyzed.

e) Bar graph showing the lineage affiliation of colonies from Lin⁻CD34⁺ normal cord blood
(CB) cells (n=5 biologically independent samples) and CD34⁺ BM cells in the CR sample of
patient #201-022. Numbers next to the bar indicate the number of colonies produced per 100
plated cells. The GM:E (granulocyte-macrophage:erythroid) ratio of colonies and the 95%
confidence interval for the GM:E ratio in normal BM are shown.

Figure 3. Enasidenib induces differentiation from an ancestral IDH2 mutant clone.

All data shown refer to samples from patient #201-023.

a) Heat map of targeted re-sequencing of mutations (rows) in single cells (columns, n=63
cells) from flow-sorted BM populations isolated pre-ENA and at CR which are shown
together. Clonal identification of each cell is shown below the heat map and the key to
mutations is denoted by letters on the right. Mutation detection key: red=detected; blue=not
detected; white= sequencing failed.

b) Clonal structure of the AML sample based on single cell genotyping (SCG). Number next
to a clone indicates the number of cells identified in that clone (data from a). The most likely
clonal structure is shown in solid arrows with alternatives in dotted arrows. (*) indicates
genotype "A" or "SIAR", which were each detected in only 1 cell. § indicates genotype "AS"
with ADO of an *IDH2* allele in 2/3 cells. Ø "SI" with ADO of the *ASXL1* allele in 3/12 cells.
See also Supplementary Fig. 12a.

c) Clonal composition in different immunophenotypic compartments pre-ENA and at CR.Number of cells studied are indicated.

d) Clonal contribution (vertical bars) to immunophenotypic stem, progenitor, myeloid precursor and terminal mature GM populations in patient samples pre-ENA and at CR (horizontal bars). Data is from SCG except for mature GM population at CR, where the flowsorted cell population was genotyped (*). Normal BM is shown for comparison of immunophenotypic populations.

e) Clonal contribution to colonies grown from CR sample (percentage of genotyped,
individually picked colonies). Key to mutations detected are as in (b). Numbers next to the
bars indicate the numbers of colonies analyzed. Lineage affiliations are as in Fig. 1d.
f) Lineage affiliation of colonies from BM CD34⁺ cells purified from CR sample compared
with normal CB (as in Fig. 2e).

Figure 4. Enasidenib induces differentiation from a terminal IDH2m clone.

All the data here are from patient #201-011.

a) Heat map of targeted re-sequencing of mutations (rows) in single cells (columns, n=110

cells) from flow-sorted BM populations pre-treatment and CR which are shown together. The
key is as in **Fig. 3a**.

b) Clonal structure of the AML sample based on SCG. The key to the panel is as in **Fig. 3b**.

* marks the genotype "DIA" where ADO was detected in 6/7 cells. Ø denotes genotype "DId".

658 No heterozygous germline SNPs were available in the *ASXL1* gene. Estimated ADO 659 frequency of the *ASXL1* allele was 12.1%. See also Supplementary Fig. 12c.

660 c) Clonal composition in different immunophenotypic compartments pre-ENA and at CR, as

661 in **Fig. 3c.**

d) Clonal contribution (vertical bars) to immunophenotypic BM haematopoietic populations in

663 patient samples pre-ENA and at CR (horizontal bars). Data is from SCG except for mature

664 GM population pre-ENA, where the flow-sorted cell population was genotyped (*).

e) Clonal contribution to colonies grown from CR sample (percentage of genotyped,
individually picked colonies) as in Fig. 3e.

f) Lineage affiliation of colonies from BM CD34⁺ cells purified from CR sample compared
with normal CB (as in Fig. 2e).

Figure 5. Mechanisms leading to relapse of enasidenib-treated patients.

a) Summary of mechanisms (rows) leading to relapse in 12 patients (columns). Selected
mutations detected at relapse by WES (all patients except #104-021) or by Heme Panel bait
capture sequencing (#104-021) are shown.

b) Longitudinal analysis of the percentage suppression of serum 2-HG concentrations prior
to enasidenib treatment (pre-ENA), at best response (CR or PR) and at relapse in 14
patients with an *IDH2* R140 codon mutation.

c-d) Serum 2-HG levels and bone marrow blast percentages prior to enasidenib treatment
(pre-ENA), at CR or CRp (complete remission without platelet recovery) during the course of
treatment (C=cycle, D=day of treatment) and at relapse in patients #201-014 (c) and #201-

679 022 (d).

e-f) Serial mutation analyses in flow-sorted blasts prior to enasidenib treatment (pre-ENA)

681 and at relapse in patients #201-014 (e) and #201-022 (f).

Figure 6. Relapse post-enasidenib occurs through clonal evolution/selection.

683

a) Patient #104-021: Heat map of targeted re-sequencing of mutations (rows) in single cells
 (n=214 cells, columns) from flow-sorted BM populations pre-treatment and at relapse which

are shown together. The key is as in **Fig. 3a**.

b) Clonal structure of patient #104-021 based on SCG. The key to the panel is as in **Fig. 3b**.

Boxes in dotted red lines highlight clones which are only detected at relapse.

c) Clonal composition in different immunophenotypic compartments pre-ENA and at relapse,

690 as in **Fig. 3c.**

d) Clonal contribution (vertical bars) to immunophenotypic BM haematopoietic populations in
 patient samples pre-ENA and at relapse (horizontal bars). Data is from SCG.

e) Patient #201-011: Heat map of targeted re-sequencing of mutations (rows) in single cells

(n=87 cells, columns) from flow-sorted BM populations at relapse. The key is as in **Fig. 3a**.

695 f) Clonal structure of patient #201-011 at relapse. * indicates 6 cells with genotype "DIA"

696 where we detected ADO in 4/5 cells in the *DNMT3A* allele. Ø denotes "DId". The estimated

ADO frequency of the ASXL1 allele was 12.1%. § indicates 6/7 "DIXF" cells where there was

ADO for the DHX15 R222G mutant allele. See also Supplementary Fig. 12c. Boxes in dotted

699 red lines as in (b).

- g) Clonal composition in different immunophenotypic compartments at relapse, as in **Fig. 3c.**
- h) Clonal contribution (vertical bars) to immunophenotypic BM haematopoietic populations in
- 702 patient samples at relapse (horizontal bars). Data is from SCG.

703 Online Materials and Methods

704 Please see also the Life Sciences Reporting Summary

705 Patient samples

706 Bone marrow (BM) or blood samples from normal donors and AML patients were obtained 707 with informed consent and collected by research ethics committee-approved Biobanks 708 (MDSBio Study, MREC 06/Q1606/110, Oxford Musculoskeletal Biobank, MREC 709 09/H0606/11: South Central Oxford C REC), Gustave Roussy (Department of Clinical 710 Hematology and Drug Development Department (DITEP), Gustave Roussy, Villejuif) and 711 MSKCC Biobanks. Cytogenetic analyses were performed in clinical laboratories. 712 Mononuclear cells (MNCs) were isolated by Ficoll density gradient. MNCs were viably frozen 713 in 90% FCS/10% DMSO in liquid nitrogen.

714 Measurement and analysis of 2-hydroxyglutarate (2-HG)

Serum samples were collected within 28 days before the first dose of enasidenib ('screening') and/or pre-dose on day 1 of each treatment cycle. 2-HG concentration was determined by liquid chromatography tandem mass spectrometry (Covance, Inc USA according to their validated method). Baseline 2-HG was either the average of the screening sample and pre-dose cycle 1 sample, or either sample if both were not available. Percentage suppression of 2-HG was determined by comparing the lowest level of 2-HG observed ontreatment to baseline.

722 Hematopoietic cell immunophenotyping

Frozen BMMNCs from normal donors or AML samples were washed with Iscove's Modified Dulbecco's Medium (Thermo Fisher Scientific, UK), 10% fetal bovine serum (Sigma, UK) and 1mg/ml bovine pancreatic DNAse I (Sigma, UK). Cells were stained for flow cytometry with antibodies in Supplementary Table 8. Analysis was carried out on either BD LSR

Fortessa or BD FACSAria Fusion (Becton Dickinson, Oxford UK). Antibodies in the lineage (Lin) depletion are: anti-CD2,CD3,CD4,CD8a,CD10,CD19,CD20 and CD235a. 7aminoactinomycin-D (7AAD, Becton Dickinson, UK) was used as a live/dead stain. Hematopoietic stem/progenitor cells were defined as subsets of Lin⁻CD34⁺ and myeloid precursor as Lin⁻CD34⁻CD117⁺ as previously described²¹. Terminally mature myeloid cells were defined as Lin⁻CD34⁻CD117⁻.

733 Mutational analysis by FoundationOne® Heme Panel

Analysis of samples in **Fig. 1a** by FoundationOne® Heme panel was conducted by Foundation Medicine, Inc. Nucleic acid libraries were prepared from DNA and RNA extracted from fresh patient BM samples and captured using custom bait-sets targeting 405 cancerrelated genes by DNA-sequencing (DNA-seq), and 265 frequently rearranged genes by RNA-sequencing (RNA-seq). Genes included in this analysis encode known or likely targets of therapies, either approved or in clinical trials, or are otherwise known drivers of oncogenesis published in the literature⁵².

741 Mutational analysis by Fluidigm Access Array

Selected samples from Fig. 1a and Supplementary Table 2 was performed using highly
 multiplexed PCR-based targeted re-sequencing with a custom panel of 373 amplicons
 covering areas of high frequency AML mutations in 35 genes, using the Fluidigm Access
 Array platform as previously described⁶.

746 Mutational analysis by whole exome sequencing (WES).

747 Genomic DNA from flow-sorted AML blasts or CD3-positive cells (germline control) was 748 purified with Allprep DNA/RNA mini or micro Kits (Qiagen, France). After exome capture with 749 SureSelect V5 Mb All Exon kit (Agilent, Les Ulis, France), paired-end 100 bp sequencing 750 was performed on HiSeq2000 (Illumina, Paris France). Read alignment to hg19 reference 751 genome was performed using the BWA algorithm, v0.7.10 with corrections using GATK 752 (v3.3.0) after removal of PCR duplicates. Variant detection was carried out with VARSCAN 753 (v2.3.7). Somatic variants (Supplementary Table 2) were selected using the following 754 criteria: minimum depth: 8x, VAF>10% in AML blasts and lower than 10% in germline

control, and p-value <0.001. All variants were manually inspected using IGV (v2.3) software.
Artifactual variants from DNA oxidation where read pair orientation bias was observed (i.e.
predominant F2R1 orientation for C to A variations or F1R2 for G to T variations) were
filtered out. Median depth at the positions reported in the **Supplementary Table 2** was 121
(range: 19-843).

760 Selection of mutations for targeted re-sequencing in populations and single cells.

761 On average 23 somatic mutations were detected per AML sample by WES. We selected 762 mutations for further validation by targeted re-sequencing and SCG based on the following 763 criteria: 1) known recurrent mutations in AML, 2) non-recurrent mutations in genes 764 commonly mutated in AML. In addition, mutations not meeting above criteria, but where the 765 VAF varied by at least 5% between sequential samples in the same patient were of interest 766 as these may be markers of clonal shifts. Where multiple mutations had similar VAFs and 767 showed similar patterns of change in VAFs in sequential samples, a representative mutation 768 was selected. This is illustrated in the example from patient #201-011 (Supplementary 769 Figure 11a). Mutations which were not validated by targeted re-sequencing were excluded 770 in subsequent data analysis. Chromosomal loss of heterozygosity detected by WES (or by 771 karyotyping) was examined using germline single nucleotide polymorphisms (SNP) present 772 in the affected chromosomal region.

773 Mutational analysis by targeted re-sequencing

774 Mutations detected by targeted re-sequencing in hematopoietic cell populations are in 775 Supplementary Table 3. Average and range of read depths for each mutation is shown in 776 Supplementary Table 4. DNA was extracted (DNeasy Blood and Tissue extraction kit, 777 #69506 Qiagen Manchester UK) from bulk and flow-sorted cells from patient samples. 778 Where material was limiting, whole genome amplification (WGA, RepliG, Qiagen, UK)) was 779 performed. Targeted PCR was performed using high Fidelity Phusion Tag polymerase (NEB, 780 UK) or KAPA2G Multiplex DNA Polymerase (KAPA Biosystems, UK) with 10ng of gDNA. 781 Primers used are in Supplementary Table 7. A second PCR reaction added Illumina 782 barcodes and sequencing oligonucleotides prior to sample purification, quantitation, pooling

and library preparation for sequencing on Illumina MiSeq (Illumina, Saffron Walden, UK).
Raw data (average depth ~996x) was aligned using Stampy (v1.0.20)⁵³. A minimum
sequencing depth of 100 was set as a threshold for inclusion of data for analysis. >94% of
reads had Phred scores of >30. VAF was obtained using the Unix 'grep' (globally search **re**gular expression and **p**rint) command line.

788 Mutant *IDH2* variant allele frequency by quantitative PCR.

789 mIDH2 VAF was assessed in gDNA extracted from flow-sorted patient blood CD14⁺ monocytes, CD16⁺ neutrophils, or polymorphonuclear neutrophils 790 (Ficoll gradient 791 purification) at various time points during enasidenib treatment. Quantitative SNP assay 792 PCR (12.5ng DNA per test) was performed using TagMan® Universal PCR Master Mix 793 (Applied Biosystems, France), TaqMan® probes (specific for either IDH2 wild type (FAM) or 794 mutated R140Q (VIC) alleles (rs121913502, Applied Biosystems, France)). PCR was 795 performed on an ABI 7500 Fast Real-Time PCR analyzer (Applied Biosystems, France) with 796 cycling conditions: initial 1min at 60°C, 10min at 95°C, followed by 40 cycles of 15sec at 797 95°C and 1min at 60°C; and then 1min at 60°C.

798 Single cell genotyping

799 Mutations interrogated by SCG are in **Supplementary Table 3**. Single cells were flow-sorted 800 into 96 well plates containing 2µl of phosphate buffered saline. WGA was carried out using 801 Single Cell RepliG kit (Qiagen, Crawley UK). Briefly, following cell lysis, alkali denaturation 802 and neutralisation, a master mix containing Phi29 polymerase, dNTPs and random 803 oligonucleotide primers was added. WGA was carried out at 30°C for 8 hours followed by 804 heat inactivation. Diluted (1:20) amplified DNA was used in single or multiplex PCR using 805 primers relevant to the sample and high Fidelity Phusion Tag polymerase (NEB, UK) or 806 KAPA2G Multiplex DNA Polymerase (KAPA Biosystems, UK). Barcoding and sequencing 807 oligonucleotides were added by PCR and sequencing performed on Illumina MiSeg 808 (Illumina, Saffron Walden, UK). ~94% of reads had Phred scores of >30. A threshold of 50 809 reads was set for analysis inclusion. VAF thresholds for determining detection of mutations 810 were determined by genotyping 48 single cells derived from normal bone marrow, and set at

the 95% confidence level (mean ± 1.96 x standard error of mean (SEM); i.e. <5% chance of

false positive, **Supplementary Table 6**).

813 Imputation of clonal structures using bulk VAFs

The most common method used to impute clonal structure is based on the assumption that the most abundant mutation resides in the earliest occurring ancestral clone. This method may be applicable to samples with linear clonal structures but it may not be able to accurately resolve more complex or branching clonal structures. Longitudinal bulk genotyping data may offer additional information, particularly where there is evidence of clonal selection or evolution.

820 Putative clonal structure is first solved for each sample independently using bulk VAF data 821 based on the rules below. In samples where there was no colony or single cell (SC) 822 genotyping, longitudinal sampling (pre-ENA, best response and relapse) can provide 823 additional information on the likely clonal structure of that patient. In the absence of colony 824 or SCG data, bulk VAFs were used to estimate the size of the most likely clones in samples. 825 In samples where there is associated colony and/ or single cell (SC) genotyping, the clonal 826 structure was re-drawn based on these data, which provide a higher confidence structure 827 with some resolution of intermediate clones. Once this clonal structure is solved, bulk VAF 828 was used to estimate the sizes of clones in samples.

Sequence of acquisition of mutations can be imputed from bulk genotyping data, using VAF as an estimate of clonal contribution (Supplementary Fig. 11b). Mutations are first ranked according to VAF: in these examples, V_A is highest, V_D is lowest.

Factors which may cause data error and bias include limited cell equivalent representation in extracted genomic DNA from small cell numbers, bias present in whole genome amplified material and PCR bias (including sequencing bias). While we were unable to control for the first two factors, we could estimate standard error of our sequencing data. We performed technical replicate genotyping of 19 unsorted BMMC populations and obtained the standard error of mean (SEM) from VAFs from each mutation. In total, 142 standard error values

were obtained, and the average SEM was 1.9% (range 0.0-20.2%), with a 95% confidence interval upper limit of 2.47%. This limits our ability to reliably distinguish between clones varying in VAF of ~2.5% (~5% of cells if mutations were heterozygous), and we were not able to impute sequence of acquisition of mutations in population genotyping with less than 2.5% difference in VAF between them. Where three or four mutations have VAF within 2.5% of each other, the average VAF of the cluster is taken as the VAF of all mutations in that cluster (Supplementary Fig.11c).

Interpreting VAFs and cellular representation in the context of loss of heterozygosity or hemizygosity.

847 We detected multiple occurrences of loss of heterozygosity (LoH, e.g. copy-loss 848 chromosomal deletions or copy-neutral uniparental disomy) and mutations, which were on 849 the X chromosome in male patients (hemizygosity). There is complexity in interpreting bulk 850 VAFs due potential mix of cells with or without LoH within a bulk population. 851 Notwithstanding this caveat, we used the simple models set out below to help interpret VAFs. 852 in different contexts. For somatic heterozygous variants (somatic mutation or germline 853 polymorphism) in autosomal chromosomes the estimated percentage cellular representation 854 is 2xVAF% (Supplementary Fig. 11d). Heterozygous variant and chromosomal deletion 855 resulting in copy-loss LoH in autosomal chromosomes is illustrated in Supplementary Fig. 856 11e. Here, there is a non-linear relationship between VAF and cellular representation 857 (Supplementary Fig. 11f). The formula we used to estimate cellular representation was: % 858 cellular representation= $[-2.777 \times (VAF ratio)^{2}]$ + (6.145 x VAF ratio) - 2.373. In 859 heterozygous variants and copy-neutral LoH (e.g. uniparental disomy) in autosomal 860 chromosomes, percentage cellular representation is 100% -(2xVAF%) (Supplementary Fig. 861 In X-linked variants in male subjects in sex chromosomes: percentage cellular 11g). 862 presentation is the same as the VAF% (Supplementary Fig. 11h).

863 Determination of clonal structures using single cell genotyping

Each single cell was assessed for detection or non-detection of mutations in that patient sample by amplicon sequencing of DNA subject to whole genome amplification. A minimum

866 coverage of 30x across an amplicons was required for a amplicon to be called. We assigned 867 the most likely sequence of acquisition of mutations based on the genotype identified in 868 cells. For example, where mutations A, B, C, D and E were identified in a sample, discrete 869 cells with genotypes A, AB, ABC, ABD, ABCD and AE may be called. In most cases the 870 sequence of acquisition, e.g. A->AB, is clear. However, the sequence of acquisition of 871 mutations during the transition for example, from AB to ABCD may not be clear due to allelic 872 drop out (ADO) i.e. the sequence of acquisition may be AB -> ABC -> ABCD or AB-> ABD-> 873 ABCD. In such cases, intermediate genotypes represented by most cells may be more likely 874 to be true. In all cases, models of clonal structures which require the least number of discrete mutational steps required are represented⁵⁴, although alternative structures, 875 876 including ones where the same mutation is acquired twice, are possible. Once the most 877 likely clonal structure is established, cells where there was failure to amplify a locus that did 878 not alter the assignment of a mutational complement (e.g. an early mutation in the hierarchy) 879 were included in the final analysis.

880 Allele Drop Out Estimation (ADO)

ADO can be measured for by two methods. First, ADO can be determined by determining the phase of germline SNPs near mutations. For patients #201-023 and #201-011, we genotyped germline SNPs which were either in-phase (i.e. on the same allele as) with a mutation or out of phase (i.e. on the opposite allele).

In-phase SNPs were rs6597996 and rs11246258, *DEAF1* N372K mutation, patient #201-011. Out-of-phase SNPs were: (i) rs4911231, *ASXL1* G646fs mutation, patient #201-023; (ii)
rs2276598, *DNMT3A* R598X, patient #201-011; (iii) rs7657364, *DHX15* R222G, patient
#201-011. The SNPs were situated between 157 bases and 4 kilobases from the mutations.
The threshold VAF for ascertaining dropout in these SNPs was <2% (homozygous reference) or >98% (homozygous variant).

In patient #201-023 (**Fig. 3a-b**) in 2/3 cells with the "AS" genotype there was ADO of one of the *IDH2* alleles (Supplementary Fig. 12a, left) and in 2/12 cells with the genotype "SI" there was ADO in the *ASXL1* allele (Supplementary Fig. 12a, right).

Our analysis also showed that in patient #201-011 *DNMT3A* R598X and R736C mutations are on different alleles and the R736C mutation is in phase with rs2276598 (Supplementary Fig. 12b).

897 By studying both VAFs of the SNP rs2276598 and the R736C mutation in cells with 898 genotype "DIA" in patient #201-011 where ADO may have occurred we determined that in 899 6/7 cells at CR, ADO of the allele that harboured the R598X mutation had occurred 900 (Supplementary Fig. 12c left). Where the SNP did not amplify, the VAF of R736C mutation 901 was informative. Similarly, in the same patient at relapse, 5/6 "DIA" cells at relapse had ADO 902 of the R598X allele (Supplementary Fig. 12c center). We also determined that in 6/7 cells 903 with the "DIXF" genotype at relapse there had been ADO of the allele harbouring the DHX15 904 R222G mutation (Supplementary Fig. 12c, right).

905 An alternate method to determine ADO more globally is to study the frequency with which a 906 variant is called homozygous (either reference or alternative) when it should be 907 heterozygote. We analysed 6 SNPs in 5 genes (ASXL1, IDH2, DNMT3A, DEAF1 and 908 DHX15) in 402 single cells from 2 patients (#201-011 and #201-023) known to be 909 heterozygous in the germline (confirmed by genotyping population of flow sorted T cells from 910 the patients). Mean VAF was 49.1% for all 6 SNPs with a near-symmetrical distribution of 911 VAFs across these single cells ranging from 0-100% (Supplementary Fig. 12d). Frequencies 912 of homozygous reference (VAF≤1%) or variant calls in the 402 cells (VAF≥99%, thresholds 913 based on analysis of known homozygous SNPs in 237 cells) were 15.9% (64 cells/402 cells) 914 and 15.2% (61 cells/402 cells) respectively (Supplementary Fig. 12d).

915 Next, if we assume that mutations in patients were heterozygous (i.e. in cases where there is 916 was no data to support uniparental disomy or copy number loss) we asked what was the 917 frequency with which mutations were called homozygous (VAF≥99%) (presumably due to

ADO). Across the 23 mutations assessed using this method, the average ADO rate of wild type allele was 10.36% (SD 5.7%, Supplementary Fig. 12e). However, we found variation in ADO rates between different patients, even at the same mutation (e.g. *IDH2* R172K), suggesting that ADO is affected by factors additional to that of the activity of the Phi21 polymerase. Where possible we have used gene-specific germline SNPs, or alternatively used sample and mutation specific ADO estimated by the 'homozygous mutant' method for our analyses.

925 Digital Droplet PCR

926 We confirmed the VAFs detected by next-generation sequencing (NGS) using digital droplet 927 PCR (ddPCR) in 17 amplicons and 113 sorted multi-cell AML populations from 5 patients 928 using the BIORAD platform as previously described²¹. There was good correlation between 929 VAF values obtained using these two methods (R²= 0.974,). There was one AML variant 930 (DHX15 R222G) where NGS gave an unexpectedly high VAF in single normal BM MNCs 931 (where the mutation was found to be absent in the bulk normal BM sample). Detection of this 932 variant in normal and AML were therefore carried out using ddPCR which showed presence 933 of the mutation in AML bulk and single cells, but confirmed its absence in normal bulk and 934 single cells.

935 Whole transcriptome sequencing (RNAseq) and analysis of alternative splicing events 936 Copy DNA libraries were prepared using extracted RNA from AML blast cells flow-sorted 937 from re- and post-relapse samples from patients #201-011 and #201-013. cDNA libraries 938 were prepared for sequencing with tagmentation and indexing using Illumina Nextera 939 Sample Preparation kit (Illumina, Saffron Walden, UK). RNA-seq data were generated as 75 bp paired-end unstranded Illumina reads. Reads were aligned using STAR(v2.4.0.1)⁵⁵ to the 940 941 human genome (GRCh37) with default parameters. On average alignment was 96.7% (range: 95.5-97.4%) with an average of 139×10^6 (range: $116\times10^6-169\times10^6$) mapped reads 942 943 per sample. Differentially spliced events (DSEs) for the wild type (pre-relapse) and post-944 relapse with spliceosome gene mutations analysed as paired samples, were identified⁵⁶ 945 using Mixture of Isoforms (MISO v0.5.4) using default parameters. An event is termed as

differentially spliced if the Bayes Factor (BF) ≥ 10 , $|\Delta PSI| > 0.2$ where PSI is 'percentage spliced in', and the event is supported by ≥ 10 reads. DSEs are classified included or skipped spliced exons (SE), alternative 3'/5' splice sites (A3SS, A5SS), mutually exclusive exons (MXE) or retained introns (RI).

950 Colony assays

25-250 cells were plated in duplicate in 1.2ml of MethoCult GFH4435 (StemCell Technologies, Manchester, UK). AML patient samples were assayed with added Enasidenib (1 μ M). The rest of the procedure was as previously described²⁰.

954 Statistical Analysis

- 955 Where applicable, statistical analyses were performed using with GraphPad Prism software 956 (v7.02) using statistical methods noted in figure and table legends.
- 957

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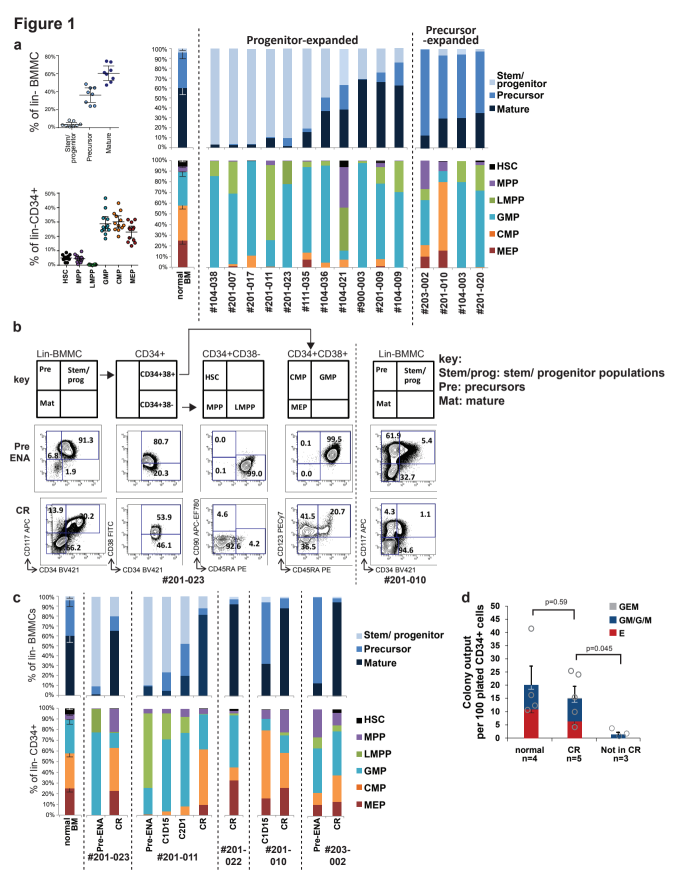
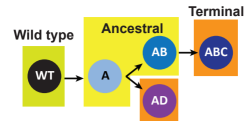


Figure 2

a Clonal origin of complete remission response to Enasidenib



b

Patient ID	#201-022	#201-023	#201-010	#201-011	#201-006	#201-027
Wild type						
Ancestral						
Terminal						

С

#201-022		IDH2 R140Q (I)	PEX26 P118S (P)	FEZ2 S208T (F)	DNMT3B N738S (D)	ZCCHC1 T712M (Z)	NPM1 L135 delins LLSIFKG (N)	ELMO3 R725Q (E)	
VAF	Pre- ENA AML blasts	36.00%	25.45%	24.96%	23.20%	22.72%	22.17%	1.83%	
	CR PBMNCs	0.24%	1.00%	1.23%	1.61%	0.58%	0.12%	0.31%	

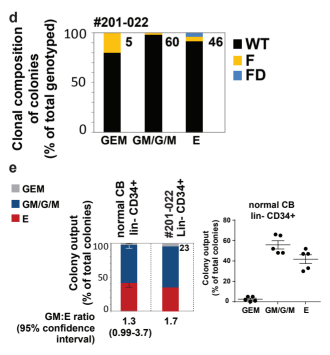


Figure 3

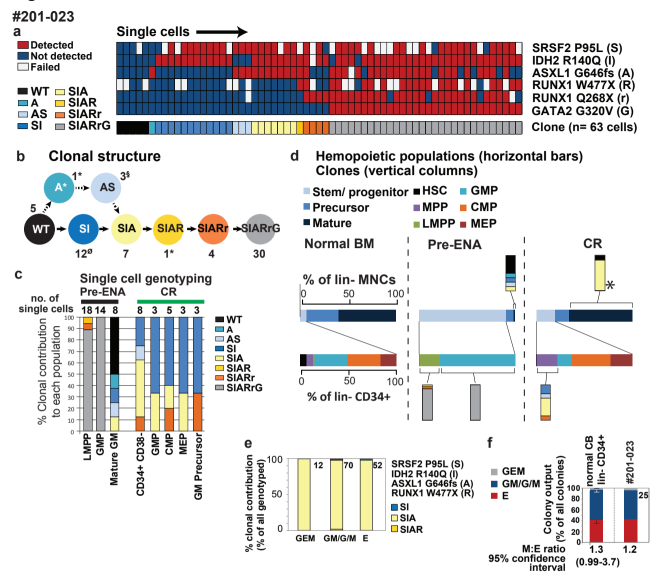


Figure 4 #201-011

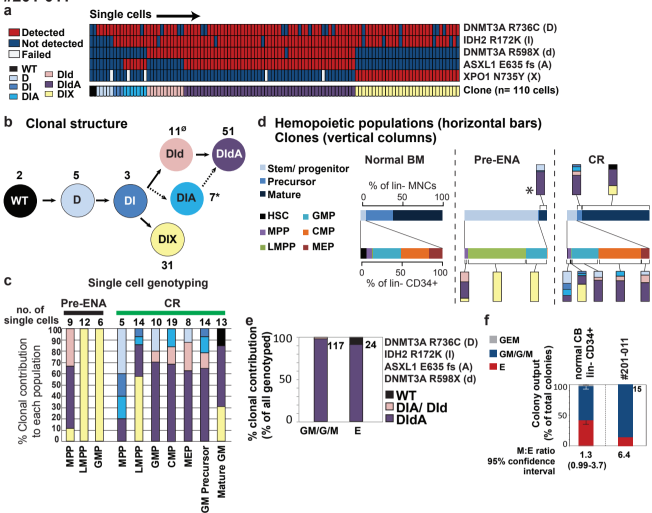
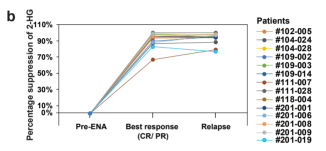
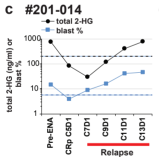


Figure 5

а

Patient ID	#201-014	#201-022	#104-021	#201-013	#201-004	#201-006	#201-010	#201-007	#201-019	#201-003	#201-011	#201-012
IDH1	IDH1 R132C	IDH1 R132H										
cytokine receptor signalling			CSF3R T618I	FLT3 D835A	FLT3 ITD			FLT3 1867M				
haematopoietic transcription factor	RUNX1 F416 Gfs*135			BCL11A N391K	GATA2 N317S/ L321R	RUNX1 T188 Hfs*25						
genes implicated in spliceosome function				DDX1 G699A							DHX15 R222G	
other genes recurrently mutated in haematopoietic cancer		ELMO3 R725Q	NFKB1 M216I	BCOR R1375W	BRCA2 A2643V				SCN3A R28H	AKAP8L D89N	DEAF1 N372K	
				CACNA1G G663R	CBLsplice acceptor				SETD1B A1054 delinsAE	PLCL1 A985V		
				UGT2B10 Y16*								
other genes recurrently mutated in other cancer				SLC18A3 W105*				MTUS Q835H		DOM3Z Y92*		
deletion chr 7q												

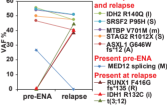




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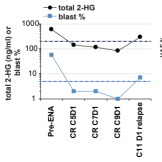
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Present pre-ENA





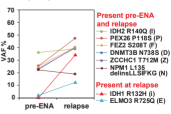


Figure 6

#104-021

