Clonal haematopoiesis is not prevalent in survivors of childhood cancer

haematopoiesis driven by leukaemia-associated Clonal somatic mutations is a common feature of ageing (Link & Walter, 2016). This phenomenon, termed clonal haematopoiesis of indeterminate potential (CHIP), is associated with an increased risk of haematological malignancies and all-cause mortality (Link & Walter, 2016). Recent empirical evidence and computational models suggest that mutation acquisition may not be the major rate-limiting factor in the emergence of CHIP (Altrock et al, 2015; McKerrell et al, 2015; Link & Walter, 2016; Young et al, 2016). Instead, clonal expansion of mutant haematopoietic stem cells (HSCs) probably reflects the interaction between the effects of driver mutations and selection pressures prevailing in the bone marrow microenvironment (Link & Walter, 2016). Notably, cytotoxic therapies have been shown to favour expansion of pre-malignant haematopoietic clones (Link & Walter, 2016). Furthermore, both adult and paediatric cancer patients treated with intensive chemoradiotherapy display an earlier onset of ageing-associated morbidities and an elevated risk of therapy-related myeloid neoplasms (t-MN) and other secondary malignancies (Rowland & Bellizzi, 2014). A recent study in adult cancer patients found that CHIP was more prevalent than in the general population and was strongly associated with t-MN and overall mortality (Gibson et al, 2017). Although CHIP is extremely rare in healthy young individuals, its prevalence and prognostic significance in paediatric cancer patients has not been studied. We therefore performed targeted deep sequencing of peripheral blood DNA from 84 childhood cancer survivors to search for subclonal mutations common in t-MN and adult clonal haematopoiesis. No individuals with somatic variants at these loci were identified. Whilst our findings could be explained by a rarity of driver mutations, the fact that human HSCs accrue somatic variants from the first decade of life (Welch et al, 2012) proposes the alternative possibility that such mutations may not confer clonal advantage in the young.

We obtained peripheral blood DNA samples from patients enrolled on long-term follow-up after treatment for a paediatric malignancy and from three age-matched controls with no cancer history. Written informed consent was obtained for sample collection and DNA sequencing from all patients or their guardian in accordance with the Declaration of Helsinki and protocols approved by the relevant institutional ethics committees (approval numbers 09REG2015, 1-09/12/ 2015). The median age at cancer diagnosis was 4-5 years, and the commonest malignancies were acute lymphoblastic leukaemia (n = 21), neuroblastoma (n = 17) and non-Hodgkin lymphoma (n = 10). Nineteen patients had received a HSC transplant (8 allogeneic and 11 autologous). The median interval between completion of cancer treatment and blood sampling was 6 years (range 2–25). Patient characteristics are summarized in Table SI.

We performed targeted next generation sequencing (NGS) using multiplex polymerase chain reaction to amplify 32 regions of 14 genes frequently mutated in CHIP or t-MN (Table I) (McKerrell et al, 2015; Link & Walter, 2016; Gibson et al, 2017). For this we extended a previously validated assay that detected clonal haemopoiesis in 2.6% of unselected adults (McKerrell et al, 2015), to include all coding exons of TP53 and PPM1D, genes implicated in t-MN pathogenesis (Rowland & Bellizzi, 2014; Link & Walter, 2016; Gibson et al, 2017). Primer design and sequencing was performed as described previously (McKerrell et al, 2015); see Table SII for primer sequences. Reads were aligned to human genome build GRCh37 using the Burrows-Wheeler Aligner (Li & Durbin, 2010) and analysed for somatic single nucleotide variants. Allele counts were generated using an in-house script (https://github.com/cancerit/alleleCount), considering only loci with ≥1000 reads and minimum base and mapping quality of 25 and 35, respectively. Somatic mutations with

Table I. Genomic regions sequenced.

Gene	Chromosome	Target codon/exon
NRAS	1	p.G12
SF3B1	2	p.K666; p.K700
DNMT3A	2	p.R882
IDH1	2	p.R132
KIT	4	exon 17
NPM1	5	exon 12
JAK2	9	p.V617
KRAS	12	p.G12
IDH2	15	p.R140; p.R172
PPM1D	17	exons 1-6
TP53	17	exons 1-12
SRSF2	17	p.P95
ASXL1	20	exon 12
U2AF1	21	p.S34; p.Q157

© 2017 The Authors. *British Journal of Haematology* published by John Wiley & Sons Ltd. doi: 10.1111/bjh.14630 This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited. variant allele frequency (VAF) \geq 0.008 (McKerrell *et al*, 2015) were sought and examined visually and by interrogation with the Shearwater algorithm (https://github.com/mg14/dee pSNV) (Gerstung *et al*, 2014).

The median sequencing depth across regions of interest was 5.3×10^3 . No somatic mutations with VAF ≥ 0.008 were observed in any of our patients or controls, demonstrating that CHIP driven by mutations at these loci is not prevalent in young individuals who have received cytotoxic treatment. By contrast, Gibson et al (2017) identified postchemotherapy CHIP (VAF > 0.02) in 29.9% of 401 adult lymphoma patients. Notably, mutations in PPM1D, a regulator of TP53, were the commonest CHIP drivers (Gibson et al, 2017). Similarly, several smaller studies have demonstrated clonal expansion in older patients undergoing chemoradiotherapy for other cancers (Link & Walter, 2016). An investigation of haematopoietic clonal dynamics in 15 adult acute myeloid leukaemia patients found that, after induction chemotherapy, five had marked expansion of clones unrelated to their leukaemia (Link & Walter, 2016). Most clones carried canonical leukaemia mutations and continued to expand years after remission (Link & Walter, 2016). In a study exploring the clonal origins of t-MN, TP53-mutated clones expanded dramatically after cytotoxic treatment, whereas the same mutations demonstrated very modest clonal advantage in healthy individuals (Link & Walter, 2016). In light of the above, our findings have two plausible explanations: (i) that somatic driver mutations are very uncommon in young individuals even after exposure to chemotherapy or (ii) that accrual of such mutations is insufficient to trigger clonal expansion in this age group. The latter is supported by findings that oncogenic mutations begin accumulating early in life (Welch et al, 2012) and that cancer-associated mutations are less able to drive clonal expansion in young compared to old stem cells (Zhu et al, 2016). The fact that bona-fide driver mutations do not always lead to haematopoietic clonal expansion, even after several years, was highlighted by Young et al (2016), using ultra-sensitive sequencing. Therefore our results should not be taken to reflect absence of potentially oncogenic HSC mutations in young cancer survivors. Rather, it is possible that even canonical leukaemogenic mutations may not commonly drive clonal outgrowth in children and young adults despite exposure to extreme haematopoietic stress, implicating age-related changes in HSCs and/or their microenvironment as key determinants of relative fitness. More sensitive DNA sequencing methods may enable detection of very rare cells harbouring known CHIP drivers mutations in similar patient cohorts, which would lend support to this hypothesis. Studies of larger numbers of paediatric cancer survivors are needed to identify rare individuals with CHIP after chemoradiotherapy, whose particular characteristics may offer insights into factors facilitating clonal outgrowth of mutated HSCs. Furthermore, in view of the shifting patterns of mutations

driving CHIP in different adult age groups (McKerrell *et al*, 2015), selective pressures particular to a less mature bone marrow environment may confer clonal advantage on a distinct spectrum of somatic variants in the very young. Although a much broader screening approach is required to identify such mutations, the potential role for CHIP as a biomarker for patient risk-stratification (Gibson *et al*, 2017) may render this a worthwhile endeavour.

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Author contributions

GSV, GC and FF conceived and designed the study. NH designed sequencing assays. GC performed experiments and bioinformatics analysis. GC and GSV wrote the manuscript with input from FF. DJ and IV wrote scripts and contributed to analysis strategy. FF, MP, MD and DC contributed to sample acquisition and patient recruitment.

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Supporting Information

Table SI. Patient characteristics**Table SII.** Primer sequences

Additional Supporting Information may be found in the online version of this article:

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