# Title: Identification of proliferative and non-proliferative subpopulations of leukemic cells in CLL

# Running title: Kinetically distinct subpopulations in CLL

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## Abstract.

Pathogenesis in chronic lymphocytic leukemia (CLL) is strongly linked to the potential for leukemic cells to migrate to and proliferate within lymph-nodes. Previous *in vivo* studies suggest that all leukemic cells participate in cycles of migration and proliferation. *In vitro* studies, however, have shown heterogeneous migration patterns.

To investigate tumor subpopulation kinetics, we performed *in vivo* isotope-labeling studies in ten patients with *IgVH*-mutated CLL (M-CLL). Using deuterium-labeled glucose, we investigated proliferation in sub-populations defined by CXCR4/CD5 and surface (sIgM) expression. Mathematical modeling was performed to test the likelihood that leukemic cells exist as distinct sub-populations or as a single population with the same proliferative capacity. Further labeling studies in two patients with M-CLL commencing idelalisib investigated the effect of B-cell receptor (BCR) antagonists on sub-population kinetics.

Modeling revealed that data were more consistent with a model comprising distinct sub-populations (p=0.008) with contrasting, characteristic kinetics. Following idelalisib therapy, similar labeling suppression across all sub-populations suggested that the most proliferative subset is the most sensitive to treatment. As the quiescent sub-population precedes treatment, selection likely explains the persistence of such residual non-proliferating populations during BCR-antagonist therapy. These findings have clinical implications for discontinuation of long-term BCR-antagonist treatment in selected patients.

**Introduction.**

The clinical course and outcome of chronic lymphocytic leukemia (CLL) varies markedly between patients. Investigation of the biological basis of this heterogeneity has contributed much to our understanding of disease pathogenesis. As for many tumors, genomic abnormalities are key determinants of outcome, however, functional properties of the tumor, especially its capacity to migrate into lymphoid tissues and proliferate in response to B-cell receptor (BCR) derived signals, are also important.

In addition to differences between patients with CLL, there is also evidence for heterogeneity within the leukemic clone within individual patients. Large scale sequencing studies have revealed significant subclonal genomic diversity 1,2 that evolves over time and in response to therapy 3. Whether this is accompanied by functional intraclonal diversity, however, remains an open question since phenotype, capacity to migrate, signal and proliferate may all change in response to anatomic location and factors in the tumor microenvironment. For example, leukemic cells within the lymph node (LN) express higher levels of the prognostic marker CD38 than those in the peripheral blood (PB) 4, whilst the chemokine receptor CXCR4 is rapidly downregulated following B-cell receptor signaling, and upregulated in its absence 5.

Intraclonal diversity has been directly studied *in vivo* using the non-radioactive isotope deuterium, administered as heavy water, to label the DNA of dividing cells in patients with CLL6,7. This technique enables the proliferation rate, lifespan and location of populations and subpopulations of tumor cells to be determined from measurements of isotope enrichment in sorted subsets over time. Labeled cells took several days to appear in the PB, compatible with proliferation within and subsequent release from lymphoid tissues, a finding later confirmed by tissue direct sampling 8. Analysis of sorted subfractions showed that the CXCR4lo/CD5hi subset appeared in the blood first and had the highest deuterium enrichment whilst CD5int/CXCR4int and CD5lo/CXCR4hi subsets show progressively lower and later labeling. In the light of these observations, it was proposed that CXCR4lo/CD5hi cells represent the most recently proliferated migrants from tissue to blood, and that, in the PB, these cells quiesce, re-express CXCR4, downregulate CD5 and then re-enter lymphoid tissue where further proliferation may occur. Since deuterium was eventually found in all sorted subsets, these data suggest that the CLL clone behaves in a homogeneous manner and that all cells are similarly able to participate in cycles of migration and proliferation.

Although this model of CLL biology, in which the leukemic clone is functionally homogeneous in respect of its capacity to migrate and proliferate, fits much of the available data, several observations indicate that function may be heterogeneous within the clone. First, although inhibitors of BCR signalling (BCRi) undoubtedly have a major impact on the size of the leukemic clone, in many patients, small numbers of leukemic cells persist 9 suggesting that not all tumor cells are equally dependent on this pathway for survival. Second, CXCR4lo/CD5hi peripheral blood CLL cells expressing a similar phenotype to LN CLL cells show preferential migration across vascular endothelium 10, suggesting the existence of a subpopulation primed for LN entry and proliferation. Finally, long term follow-up of patients who have undergone whole exome sequencing has shown a clear relationship between subclonal complexity and tumor growth characteristics *in vivo* 11 .

In view of the above, we re-examined whether the leukemic clone in CLL is always homogeneous in respect of proliferative potential. The deuterium labeling studies described above used heavy water (2H20) which, being administered over several weeks, labels multiple generations of cells. Such studies therefore have a limited capacity to resolve subpopulation evolution over time. To address this issue, we opted to use deuterated-glucose (2H2-glucose) as a labeling agent. Because it has a very short half-life it labels a cohort of cells dividing over a period of less than a day and allows the progress of that population of cells to be tracked through subsequent phenotypic transitions and migration. Although CLL cells divide slowly, the high precursor enrichments achieved with deuterated-glucose (about one log higher than 2H2O) allow measurable labeling after a short-pulse label. We have previously used this approach to estimate the lifespan of CLL cells, generating average results very similar to those estimated using 2H2O 12.

In the present study, we focused our investigation on patients with M-CLL, the subgroup of patients most likely to exhibit persistent lymphocytosis during BCR inhibitor therapy 9, and therefore most likely to have more readily detectable functional subclonal heterogeneity. Mathematical modeling techniques were then used to determine whether subsets defined by phenotypic markers associated with migration and BCR signalling capacity (CXCR4/CD5, surface IgM 13 and BCR internalization 14) are more likely to be homogenous, deriving from each other (Model A, Figure 1), or instead behave as separate subpopulations that remain distinct over time (Model B, Figure 1). We also report the impact of the BCR antagonist idelalisib on the release and survival of CLL subpopulations labeled prior to and again during therapy.

## Methods.

### Patients and Sampling.

Two cohorts of subjects with CLL with mutated immunoglobulin heavy chain genes (*IGHV*) were studied (Table 1). In the first, 10 treatment-naïve patients with non-progressive Binet Stage A or B M-CLL were labeled with oral deuterium-glucose; PB was drawn and CLL cells isolated at set intervals after labeling (Figure 2). Duplicate studies were performed 8 weeks later to estimate reproducibility.

The second cohort comprised two individuals participating in the CALiBRe study (EudraCT 2012-003631-36) which investigated idelalisib, a phosphoinositide-3 kinase (PI-3K) inhibitor, in patients with previously-untreated symptomatic CLL15; deuterium labeling was included as an optional sub-study. CALiBRe closed early due to safety concerns with idelalisib, hence recruitment to the labeling sub-study was limited to two patients. In these subjects, deuterium-glucose administration was timed to begin 24 hours before commencing idelalisib treatment (150 mg bd); labeling and sampling matched those in the untreated cohort (Figure 2). All subjects gave written, informed consent; studies were approved by the UK National Research Ethics Service (13/LO/0434 and 15/YH/0020).

**Deuterium-glucose labeling and analysis.**

*In vivo* labeling was performed by oral administration of 60g deuterium-glucose (6,6-2H2-glucose, Cambridge Isotope Laboratories, USA) over 10 hours, as previously described 16. Since labeled glucose disappears rapidly (within hours), this approach labels cells dividing during an approximately 12-hour window. PBMC from follow-up samples (Figure 2) were cryopreserved in 10% dimethyl sulphoxide prior to sorting (as below), then analysed for DNA deuterium enrichment by gas chromatography-mass spectrometry (GC-MS) as previously described 16,17. Results were expressed as the percentage of newly-proliferated cells per day after normalizing for the level and duration of blood glucose labeling directly measured during the labeling phase, as previously described 16.

### Immunofluorescence Cell Sorting.

Cryopreserved cells were thawed and stained with fluorochrome-conjugated antibodies to CD5/CD19 either alone or in combination with either CXCR4, sIgM or IgM linked to pHrodo, a dye that fluoresces following internalization into acidic endosomes; for details see Supplementary Table 1. Cells were then flow sorted into fractions representing the whole CLL clone (CD5+/CD19+) or subpopulations defined by expression of CD5 and CXCR4, or IgM at the cell surface (sIgM) or within acidified endosomes (eIgM). Each selected CLL subpopulation comprised 10-15% of the whole CLL cell population, which appeared as a continuum (Figures 4a, 5a and 5c). Fluorochromes and gating strategies are detailed in Supplementary Methods.

### Statistical analysis and Mathematical Modeling.

We fitted mathematical models describing the kinetics of CLL cells to the glucose enrichment data. The models assumed that CLL cells undergo proliferation in a compartment “tissue” that is not in immediate dynamic equilibrium with blood, i.e. we assumed that cells neither divide in blood nor in a compartment that freely and immediately recirculates with blood; this assumption precludes a model in which PB1 immediately recirculates with “tissue”. The precise anatomical correlate is immaterial to the modeling. In addition to tissue we included three PB subsets: PB1, PB2, and PB3 representing the three sorted subpopulations: PB1: CXCR4lo/CD5hi, sIgMhi or eIgMhi; PB2: CXCR4int/CD5int,sIgMint or eIgMint; PB3: CXCR4hi/CD5lo, sIgMlo or eIgMlo. Since sIgM and eIgM levels in CLL are closely correlated14, for the purposes of this study the two were considered together.

Two models of CLL kinetics were considered. In the first (Model A) there are no distinct subpopulations; instead, newly-proliferated CLL cells transit from the tissue compartment, through pools PB1, PB2 and PB3 then back to tissues. In the second (Model B), CLL cells transit from tissues to blood pools PB1, then PB2, then back to tissues, whilst the third subpopulation, PB3 (CXCR4hi/CD5lo, sIgMlo or eIgMlo cells), does not re-enter tissues but forms a stable non-proliferating subpopulation in blood. Both models were fitted to deuterium labeling data from bulk and sorted subsets. The small sample corrected Akaike Information Criterion (AICc) was used to determine which model best described the data (based on fit to data penalized by model complexity). Model parameters are fully defined and equations describing the movement between tissue and blood compartments provided in Supplementary Methods.

## Results.

### Proliferation, release, and disappearance rates of labeled CLL cells.

DNA deuterium labeling curves for bulk tumor cells from this cohort of patients with non-progressive M-CLL are shown in Figure 3 (for detail of days 0-7 see Supplementary Figure 2). The curves reflect three processes; (i) the first part of the curve, generally a delay before the appearance of labeled cells, represents the time taken for cells to transit between their site of division and blood, the site of sampling; (ii) the peak primarily reflects the proliferation rate – more proliferation resulting in greater uptake of deuterium during the short labeling phase; (iii) the subsequent disappearance of label from blood primarily reflects the rate at which labeled cells die or migrate out of blood. Analysis of DNA deuterium enrichment curves with these processes in mind led us to four general observations.

Firstly, we noted striking inter-individual heterogeneity in labeling patterns, particularly evident in the release rates of newly proliferated CLL cells into blood. While some subjects such as K8 and K10, demonstrated rapid release of highly-labeled cells, others, such as K5, K6, K9, showed slower release with a lower peak, and some (e.g. K1) showed very little labeling at all. The time to maximum PB deuterium enrichment ranged from 2 to 56 days (median 10.5 days) and was longest in patients with splenomegaly; four of the five patients with very delayed release (≥14 days) had splenomegaly (K4, K5, K6, K9).

Secondly, we observed generally low deuterium labeling rates in tumor cells, consistent with previously reported low proliferation rates in CLL cells 12,18,19. In this study, taking peak labeling as a crude minimum estimate of proliferation (disregarding cell death), we estimated a median proliferation rate of 0.26 %/d (IQR = 0.20-0.32 using cycle 1 data, n=10) equivalent to a median half-life of 271 days (0.29%/d if also including cycle 2 data, n=19).

Thirdly, we found that labeled CLL cells disappeared remarkably slowly (Figure 3); labeled cells remaining detectable at 56 days in 5/10 cases, suggesting that death rates of labeled cells are low (i.e. that CLL cells have a long lifespan), in keeping with previous observations6. Fourthly, although there was wide inter-patient variability, intra-patient reproducibility in repeat studies was generally good (Supplementary Figure 3).

**CXCR4/CD5 expression predicts proliferative subclonal behaviour and identifies a non-proliferative subpopulation.**

### Analysis of subfractions of PB CLL cells sorted into three non-contiguous populations according to CXCR4 and CD5 expression (Figure 4) showed that the CXCR4lo/CD5hi subfraction appeared in the PB earliest, consistent with previous studies7. The intermediate population (CXCR4int/CD5int) contained less deuterium and took longer to appear in PB, compatible with progressive acquisition of CXCR4 and loss of CD5 expression by the labeled cohort of cells (Figure 4). The CXCR4hi/CD5lo subpopulation, however, exhibited minimal or undetectable labeling throughout the eight-week period of monitoring. Patient K5 proved to have two phenotypically distinct clones and was not included in subsequent analyses.

### Expression of BCR identifies subclonal populations with distinct kinetic characteristics.

In view of the previous observation that expression of sIgM is dynamic and that low expression is related to a reversible state of anergy, we wenton to investigate proliferation in CLL subsets defined by sIgM expression

5,13,20 (Figure 5a). Assessment of deuterium labeling revealed that CLL cells with the highest sIgM expression (sIgMhi) represent the most recently proliferated population with progressively lower labeling in sIgMint and sIgMlo subpopulations (Figure 5b). Interestingly, in some patients, no detectable label appeared in the sIgMlo cells over the entire two months of the study. We then went on to further investigate the functional relationship between subpopulations defined by the presence of IgM within acidified endosomes (eIgM) in three patients; this measure reflects BCR internalisation and correlates with surface expression14. We found that cells with high rates of BCR internalisation (eIgMhi) had the highest rates of proliferation whilst the subpopulation with the lowest levels of eIgM remained unlabeled suggesting a fixed state of quiescence (Figure 5c&d).

### Mathematical modeling frequently predicts the presence of independent subpopulations.

We next assessed the relationship between subpopulations using mathematical modeling, testing two alternative hypotheses: (i) that low labeling in the CXCR4hi/CD5lo, sIgMlo, eIgMlo populations could be explained by dilution in sequentially labeled pools, or (ii) that CXCR4hi/CD5lo, sIgMlo, eIgMlo cells represent a distinct quiescent population. These hypotheses are represented by the two models A and B respectively (Figure 1), to which the deuterium labeling data was fitted.

Overall, across all data sets (i.e. pooling datasets sorted by CXCR4/CD5 with those sorted by sIgM and eIgM, n=14) Model B provided a substantially better description of the data (difference in AICc >1) in 7 analyses, whereas Model A only outperformed Model B in one case (Table 2). Of these 7 cases, the difference in AICc was large (>=3) for 5 datasets (Table 2). In the other 6 cases, the AICc did not discriminate between the models. Model fits are shown in Supplementary Material. We were therefore able to reject the null hypothesis that Model A always wins with certainty (p=0.008 Wilcoxon signed rank test two sided). This demonstrates that it is unlikely that there is a continuous single population of leukemic cells and that, at least in some patients, recirculation to LNs most likely occurs at the intermediate stage (Model B).

### Idelalisib has a heterogeneous effect on functional subclones.

To address the effect of BCR inhibition on these functionally heterogeneous subpopulations, we performed *in vivo* labeling in two patients (CAL01 and CAL02) with symptomatic CLL participating in the CALiBRe clinical trial. Both had mutated *IGHV* genes and CD38 expression rates >7% (Table 1). Total lymphocyte count and expression of sIgM by CLL cells was recorded over time to assess the kinetics and phenotype of the leukemic clone following drug administration. As anticipated, there was a rapid rise in the lymphocyte count in the days following treatment initiation, peaking on approximately day 4 and falling thereafter (Figure 6a). This correlated with an abrupt rise in sIgM expression, again maximal on day 2-4 before returning to baseline by day 7, falling to a mean of 43% of peak values by day 56 (Figure 6b).

Labeling with deuterium-glucose started 24 hours before the commencement of idelalisib and was repeated eight-weeks later. The first cycle of pre-treatment labeling was planned to label all cells actively dividing *in situ* in the absence of drug; idelalisib treatment would not therefore be expected to affect labeling rates in tissue as any deuterium-labeled glucose would have cleared well before idelalisib administration. According to the previous experiments in untreated patients above, such labeled cells would normally remain at their sites of replication for several days before sequential release into blood, resulting in a peak in PB labeling at ~7-14 days (Figure 4). By contrast, both study patients showed a prompt increase in labeling rates in PB CLL cells, apparent within four hours of idelalisib dosing (Figure 6 and in detail for days 0-7 in Supplementary Figure 4). Labeling peaked on day 4 and was greatest in the sIgMhi compartment; by contrast, labeling in the sIgMlo compartment remained undetectable indicating that sIgMhi cells represent the most recently proliferated population.

Strikingly, when rates of labeling were re-evaluated eight weeks into idelalisib treatment, there was a dramatic, near-complete cessation of proliferation in all leukemic cell subsets (Figure 6c), evident as the absence of an expected second peak in deuterium enrichment. In untreated individuals this peak occurred predictably 0-7 days after labeling (Figure 5b), especially in the sIgMhi subpopulation and, to a lesser extent, in sIgMint cells; for the second (d56) labeling, this peak would be expected at 56-63 days post-study initiation. Comparing the red lines at d0-7 in Fig 5b with those at d56-63 in Figure 6c, the extent to which proliferation appears to be blocked can be appreciated. At this time-point, both treated patients still had a persisting lymphocytosis but median sIgM expression levels were reduced compared to pre-treatment levels and labeling rates were now virtually undetectable in all sorted subpopulations suggesting that CLL cells expressing high levels of sIgM are most sensitive to BCR-directed treatment.

## Discussion.

In this study, we investigated whether the leukemic cells of CLL patients are homogeneous in their capacity to undergo proliferation or contain distinct subpopulations that differ in this respect. Our results convincingly show that CLL does not always behave as though the tumor is functionally homogeneous and that, at least in this population of patients with mutated *IGHV*, distinct subsets of cells with different proliferative capacities can be identified using phenotypic markers.

Uniquely, our study used *in vivo* pulse-chase labeling with deuterated-glucose to investigate this important question. To increase our capacity to detect non-proliferating subpopulations, we focussed on patients with mutated *IGHV* who are already known to have lower proliferation rates*.* As expected, we found very low estimated rates of *in vivo* tumor proliferation (median 0.26%/day), consistent with previous reports with deuterated glucose (0.57 %/d)12 or deuterium-labeled water (0.36 %/d)19, albeit in different patient populations. The delay to peak appearance of newly-produced leukemic cells in blood is consistent with cell division in the non-blood compartment for all subclones. This lag was highly variable and longest in the four patients with marked splenomegaly, suggesting that spleen and blood together form recirculating pool; when this is large, as in splenomegaly, labeling is slow. In keeping with previous studies, we found that the rates of disappearance of CLL cells from the peripheral blood were much slower than those reported for healthy B lymphocytes 12.This finding might result from impaired apoptosis in CLL cells 12, but could also reflect trapping of CLL cells in blood.

Analysis of the deuterium content of flow-sorted subpopulations clearly revealed sequential incorporation into CXCR4lo/CD5hi, then CXCR4int/CD5int, then CXCR4hi/CD5lo fractions, consistent with a model in which CXCR4lo/CD5hi cells represent the actively-dividing population. (The alternative hypothesis that all subpopulations divide but appear sequentially because they have different release dynamics is refuted by the observation that CXCR4lo/CD5hi cells comprise the population enriched for proliferating cells within the LN compartment 8.) However, our observations that deuterium enrichments were substantially lower in CXCR4int/CD5int cells, and even lower (occasionally undetectable) in the CXCR4hi/CD5lo fraction, suggests that exchange of deuterium between phenotypic groups over time may be limited, in contrast to the conclusions of previous work using deuterated water7.

Similar patterns were observed when cells were sorted according to sIgM or eIgM levels, both markers of BCR signaling capacity. Deuterium labeling was greatest and most rapid in the sIgMhi/ eIgMhi fraction, consistent with high sIgM expression in LNs, the sites of antigen encounter and proliferation 14,21. (Although it may seem counterintuitive that cells remain sIgMhi despite recent BCR ligation, levels may be maintained by microenvironment-related factors such as CD4 T-cell derived IL4 which enhances sIgM expression and BCR signaling whilst inducing reciprocal downregulation of CXCR4 22,23.) The most striking finding, however, was that, as for CXCR4/CD5 sorting, a subpopulation defined by reduced surface or endosomal BCR expression remained almost entirely unlabeled throughout the period of the study.

When these patterns of labeling of CLL subpopulations were analysed using mathematical modeling we found very little support for sequential labeling model (Model A) and refuted the null hypothesis that ‘Model A is true in all cases of M-CLL’. Conversely, our data suggest that, at least in some if not all patients, most CLL cells return to LN as CXCR4int/CD5int / sIgMint, eIgMint cells, to re-emerge after proliferation as CXCR4lo/CD5hi/sIgMhi/eIgMhi cells. Meanwhile, the CXCR4hi/CD5lo/sIgMlo/eIgMlo subpopulation forms a quiescent long-lived pool of cells that either does not migrate and proliferate, or if it does, only at rate lower than the limit of detection of the method described, a possibility which is perhaps more likely as some input of new cells is required to balance cell death and preserve population size. In terms of phenotypic characterisation, previous studies have shown that the CXCR4hi/CD5lo population is enriched for CCR7 and CTLA4 expression, in contrast to the CXCR4lo/CD5 fraction which is enriched for CD38 and CD62L7.

A hypothetical model, building upon previously-published models 7 is shown in Figure 7. This interpretation is consistent with work performed using an *in vitro* circulation system showing that the leukemic clone has heterogeneous kinetics and that migration is limited to a small sub-population characterised by an activated lymph-node phenotype 10,24. It is also consistent with recent studies showing that subclonal diversity correlates with the *in vivo* growth characteristics of CLL11. Interestingly, in their study, logistic tumor growth pattern, with plateauing of cell numbers after an initial increase, was observed in patients with lower subclonal complexity and mutated *IGHV*, and it is plausible that this may be a consequence of the emergence of the non-proliferating subset described herein.

Finally, although necessarily limited to two patients by regulatory issues beyond our control, pre-treatment deuterium labeling showed that idelalisib therapy induces rapid release of recently-proliferated sIgMhi cells into PB, consistent with the observation that Ki67

25 and sIgM expression 26 in circulating CLL cells rapidly rises following initiation of treatment with ibrutinib, likely as a consequence of release from lymphoid tissues. After eight weeks of treatment with idelalisib, deuterium labeling data showed cessation of proliferation of the sIgMhi subpopulation, as expected from reports that CLL cells expressing high levels of sIgM (the main determinant of BCR signaling) are most sensitive to BCR inhibition 13,20. The quiescent sIgMlo population meanwhile persisted, perhaps because it is less dependent on BCR signaling and so less sensitive to its inhibition (Supplementary Figure 6). Selection of the non-proliferating CXCR4hiCD5lo, sIgMlo subset may explain the persistent low-level disease observed during long term ibrutinib therapy 9. This idea is supported by changes in the tumor immunophenotype over time during ibrutinib therapy as sIgM, eIgM and CD5 expression levels initially increase then fall in the longer term 14,26.

In terms of limitations, although we studied only 10+2 subjects, our conclusions are supported by replicate measurements for every sample/time-point and repeated measurements over time within subjects, giving an overall database of over 500 enrichment measurements (Supplementary Material). Inter-individual variation was marked, as would be expected in such a heterogenous disease, and the overall pattern of labelling varied widely between patients (Figure 3), but despite this the hierarchy and sequential nature of labeling in defined subpopulations was remarkably similar between patients (Figures 4 & 5). Our primary finding regarding model comparisons is highly unlikely to have arisen by chance (p=0.008).

Our results raise important further questions. Firstly, how do these functionally distinct subsets arise? The link between anergy and lack of capacity to migrate into LN follicles has been previously described in normal murine B-cells 27. Our hypothesis is that the non-migratory/non-proliferating CLL subset we have identified resembles anergic normal B-cells in that both are excluded from entry into follicles but, whereas the latter undergo apoptosis, CLL cells are rescued by *bcl-2* expression. Detailed genetic and biochemical studies were outside the scope of the present study, but would determine whether the previously identified clonal heterogeneity 1,11,28 is linked to the functional and kinetic differences we have described here. Secondly, if there is a non-proliferative component in some cases of CLL, is it necessary to eliminate every detectable leukemic cell to obtain lasting clinical benefit? Many patients with monoclonal B-lymphocytosis or stage A CLL remain well for prolonged periods; if it is possible to eliminate the more dangerous proliferative component of the disease with drugs such as ibrutinib, then perhaps therapy can be discontinued once only non-proliferative/anergic cells remain. The observation of prolonged disease stability in some patients who have discontinued ibrutinib because of side effects suggests that this may be the case 29.

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## Competing interest disclosure.

## None of the authors has a relevant conflict of interest

## Author Contribution Statement.

K.C designed the research, performed the research, analyzed data and wrote the paper.

Y.Z performed research

E.C performed research

L.B performed research

A.P designed research, analyzed data and wrote the paper

B.A\* designed research, analyzed data and wrote the paper

S.D\* designed research, analyzed data and wrote the paper

D.M\* designed research, analyzed data and wrote the paper

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**References.**

1. Schuh A, Becq J, Humphray S, Adrian A, Burns A, Clifford R et al. Monitoring chronic lymphocytic leukemia progression by whole genome sequencing reveals heterogeneous clonal evolution patterns. *Blood*. 2012;120(20):4191-4196.

2. Wu CJ. CLL clonal heterogeneity: an ecology of competing subpopulations. *Blood*. 2012;120(20):4117-4118. d

3. Landau DA, Carter SL, Stojanov P, McKenna A, Stevenson K, Lawrence M et al. Evolution and Impact of Subclonal Mutations in Chronic Lymphocytic Leukemia. *Cell*. 2013;152(4):714-726.

4. Patten PEM, Buggins AGS, Richards J, Wotherspoon A, Salisbury J, Mufti G et al. CD38 expression in chronic lymphocytic leukemia is regulated by the tumor microenvironment. *Blood*. 2008;111(10):5173-5181.

5. Vlad A, Deglesne PA, Letestu R, et al. Down-regulation of CXCR4 and CD62L in Chronic Lymphocytic Leukemia Cells Is Triggered by B-Cell Receptor Ligation and Associated with Progressive Disease. *Cancer Research*. 2009;69(16):6387-6395. d

6. Calissano C, Damle RN, Hayes G, Murphy E, Hellerstein M, Moreno C et al. In vivo intraclonal and interclonal kinetic heterogeneity in B-cell chronic lymphocytic leukemia. *Blood*. 2009;114(23):4832-4842.

7. Calissano C, Damle RN, Marsilio S, Yan X, Yanocopoulos S, Hayes G et al. Intraclonal Complexity in Chronic Lymphocytic Leukemia: Fractions Enriched in Recently Born/Divided and Older/Quiescent Cells. *Mol Med*. 2011;17(11-12):1374-1382.

8. Herndon TM, Chen S-S, Saba NS, Valdez J, Emson C, Gatmaitan M et al. Direct in vivo evidence for increased proliferation of CLL cells in lymph nodes compared to bone marrow and peripheral blood. *Leukemia*. 2017;31(6):1340-1347.

9. Woyach JA, Smucker K, Smith LL, Lozanski A, Zhong Y, Ruppert A et al. Prolonged lymphocytosis during ibrutinib therapy is associated with distinct molecular characteristics and does not indicate a suboptimal response to therapy. *Blood*. 2014;123(12):1810-1817.

10. Pasikowska M, Walsby E, Apollonio B, Cuthill K, Phillips E, Coulter E et al. Phenotype and immune function of lymph node and peripheral blood CLL cells are linked to transendothelial migration. *Blood*. 2016;128(4):563-573.

11. Gruber M, Bozic I, Leshchiner I, Livitz D, Stevenson K, Rassenti L et al. Growth dynamics in naturally progressing chronic lymphocytic leukaemia. *Nature*. June 2019:1-20.

12. Defoiche J, Debacq C, Asquith B, et al. Reduction of B cell turnover in chronic lymphocytic leukaemia. *Br J Haematol*. 2008;143(2):240-247.

13. Ian Mockridge C, Potter KN, Wheatley I, Neville LA, Packham G, Stevenson FK. Reversible anergy of sIgM-mediated signaling in the two subsets of CLL defined by VH-gene mutational status. *Blood*. 2007;109(10):4424-4431.

14. Coulter EM, Pepper A, Mele S, Folarin N, Townsend W, Cuthill K et al. In vitroand in vivoevidence for uncoupling of B-cell receptor internalization and signaling in chronic lymphocytic leukemia. *Haematologica*. 2018;103(3):497-505.

15. Hallek M, Cheson BD, Catovsky D, Caligaris-Cappio F, Dighiero G, Dohner H et al. Guidelines for the diagnosis and treatment of chronic lymphocytic leukemia: a report from the International Workshop on Chronic Lymphocytic Leukemia updating the National Cancer Institute–Working Group 1996 guidelines. *Blood*. 2008;111(12):5446-5456.

16. Macallan DC, Asquith B, Zhang Y, de Lara C, Ghattas H, Defoiche J et al. Measurement of proliferation and disappearance of rapid turnover cell populations in human studies using deuterium-labeled glucose. *Nat Protoc*. 2009;4(9):1313-1327.

17. Busch R, Neese RA, Awada M, Hayes GM, Hellerstein MK. Measurement of cell proliferation by heavy water labeling. *Nat Protoc*. 2007;2(12):3045-3057.

18. Burger JA, Li KW, Keating MJ, Sivina M, Amer A, Garg N et al. Leukemia cell proliferation and death in chronic lymphocytic leukemia patients on therapy with the BTK inhibitor ibrutinib. *JCI Insight*. 2017;2(2):e89904.

19. Messmer BT, Messmer D, Allen SL, Kolitz J, Kudalkar P, Cesar D et al. In vivo measurements document the dynamic cellular kinetics of chronic lymphocytic leukemia B cells. *J Clin Invest*. 2005;115(3):755-764.

20. Coelho V, Krysov S, Steele A, Hidalgo M, Johnson P, Chana P et al. Identification in CLL of circulating intraclonal subgroups with varying B-cell receptor expression and function. *Blood*. 2013;122(15):2664-2672.

21. Herishanu Y, Pérez-Galán P, Liu D, Biancotto A, Pittaluga S, Vire B et al. The lymph node microenvironment promotes B-cell receptor signaling, NF-kappaB activation, and tumor proliferation in chronic lymphocytic leukemia. *Blood*. 2011;117(2):563-574.

22. Guo B, Zhang L, Chiorazzi N, Rothstein TL. IL-4 rescues surface IgM expression in chronic lymphocytic leukemia. *Blood*. 2016;128(4):553-562.

23. Aguilar-Hernandez MM, Blunt MD, Dobson R, et al. IL-4 enhances expression and function of surface IgM in CLL cells. *Blood*. 2016;127(24):3015-3025.

24. Walsby E, Buggins A, Devereux S, Jones C, Pratt G, Brennan P et al. Development and characterization of a physiologically relevant model of lymphocyte migration in chronic lymphocytic leukemia. *Blood*. 2014;123(23):3607-3617.

25. Herman SEM, Niemann CU, Farooqui M, Jones J, Mustafa R, Lipsky A et al. Ibrutinib-induced lymphocytosis in patients with chronic lymphocytic leukemia: correlative analyses from a phase II study. April 2014:1-9.

26. Drennan S, Chiodin G, D'Avola A, Tracy I, Johnson P, Trenton L et al. Ibrutinib Therapy Releases Leukemic Surface IgM from Antigen Drive in Chronic Lymphocytic Leukemia Patients. *Clinical Cancer Research*. 2019;25(8):2503-2512.

27. Cyster JG, Hartley SB, Goodnow CC. Competition for follicular niches excludes self-reactive cells from the recirculating B-cell repertoire. *Nature*. 1994;371(6496):389-395.

28. Landau DA, Wu CJ. Chronic lymphocytic leukemia: molecular heterogeneity revealed by high-throughput genomics. *Genome Med*. 2013;5(5):47.

29. Shanafelt TD, Wang V, Kay NE, Hanson C, O'Brien S, Barrientos J et al. Ibrutinib and Rituximab Provides Superior Clinical Outcome Compared to FCR in Younger Patients with Chronic Lymphocytic Leukemia (CLL): Extended Follow-up from the E1912 Trial. *Blood*. 2019;134(Supplement\_1):33-33.

**Figure Legends**

**Figure 1. Models for CLL cell proliferation and recirculation**

(a) Model A represents the hypothesis that there is a single recirculating pool of cells. We assume all proliferation (p) occurs in a non-blood compartment, denoted “Tissue” and that the blood compartment can be considered as 3 phenotypically distinct subpopulations. Black arrows denote transition between compartments and grey arrows denote cell death or disappearance. (b) Model B represents the alternative hypothesis that there are two (semi-)independent cell populations. The PB2 compartment recirculates to the tissue compartment, leaving PB3 as a distinct non-recirculating pool of cells. Cell disappearance (db, disappearance from blood compartment; dl, disappearance from non-blood compartment is thought (but not assumed) to be primarily due to cell death. Version with all parameters marked is shown in Supplementary Figure 1.

**Figure 2. Study Schemas**

In the observational study, 10 patients underwent labeling with deuterium-labeled glucose (6,6-2H2-glucose), shown by the yellow bars, on two occasions (Cycles 1 and 2) separated by eight weeks, denoted day 0 and day 56 respectively. In the CALiBRe study, the labeling schedule was performed in an identical way but treatment with idelalisib was commenced on day 1, starting 24 hours after commencement of the first labeling cycle, and continued for a minimum of 16 weeks (as indicated by the red arrow). Blood was drawn at regular intervals as shown by the blue arrows (Cycle 1: baseline, daily for the first 4 days, then weekly for 4 weeks; Cycle 2: baseline, day 4, weekly for 4 weeks, then on day 56).

**Figure 3. Deuterium labeling in CLL cells in ten patients with non-progressive disease**

DNA deuterium enrichments in purified CLL cells from 10 subjects (K1-K10) without intervention. Data are expressed as fraction of new cells produced (F, as a percentage) normalised by the glucose enrichment area under curve to the equivalent of one day’s labeling, i.e equivalent to %/day. Note different axes for K8 and K10. Cycle 2 results were similar (Supplementary Figure 2). Time represents days following oral labeling with deuterium-labeled glucose (labeling day = d0); error bars represent the standard deviation of at least four replicate measurements by GC-MS (error bars not shown where they are smaller than the symbols).

**Figure 4. Deuterium labeling for CLL subpopulations defined by CXCR4 and CD5 expression**

(a) Sorting protocol for CLL cells according to reciprocal and intermediate expression of CXCR4 and CD5. Gates were set to collect 15% of events within each subpopulation: CXCR4lo/ CD5hi (red arrow), CXCR4int/CD5int (blue arrow), and CXCR4hi/CD5lo (green arrow). (b) Deuterium enrichment curves of sorted CLL cells: CXCR4loCD5hi (red circles), CXCR4intCD5int (blue squares), and CXCR4hiCD5lo (green triangles). Results are shown from 7 subjects (K2, K3, K4, K5, K6, K9, and K10), expressed as fraction of new cells (F %) normalised to one day’s labeling; error bars represent the standard deviation of ≥4 replicate measurements by GC-MS. The F scale varies between subjects to best represent deuterium enrichment. Time represents days post-labeling. Note that K5 had a dual population of CLL cells.

**Figure 5. Association of CLL kinetics with surface IgM (sIgM) expression and expression within acidified endosomes (eIgM)**

(a) Sorting protocol for CLL cells according to sIgM expression. Gates were set to include 10% of events within each subpopulation: sIgM hi (red arrow), sIgM int (blue arrow), and sIgM lo (green arrow). (b) Deuterium enrichment curves of sorted CLL cells: sIgMhi (red circles), sIgM int (blue squares), sIgMlo (green triangles). Results are shown from 6 subjects (K4, K6, K7, K8, K9 and K10). (c) Sorting protocol according to expression of BCR within the acidified endosome (eIgM). Gates were set to include 10% of events within each subpopulation: eIgM hi (red arrow), eIgM int (blue arrow), and eIgM lo (green arrow). (d) Deuterium enrichment curves of sorted CLL cells: eIgMhi (red circles), eIgM int (blue squares), and eIgMlo (green triangles). Results are shown from 3 subjects (K6, K9 and K10). For (b) and (d) data are expressed as fraction of new cells (F %) normalised to one day’s labeling; error bars represent the standard deviation of ≥4 replicate measurements by GC-MS. F Scales vary between subjects to best represent deuterium enrichment. Time represents days post-labeling.

**Figure 6. Kinetics of lymphocytosis, median sIgM expression and deuterium enrichment in CLL cells following treatment with idelalisib (CALiBRe Study)**

Results are shown for 2 subjects: CAL01, left panels, and CAL02, right panels, pre- (day 0) and up to 84 days post-treatment with idelalisib: (a) Total lymphocyte count; (b) Median expression of sIgM; (c) Deuterium enrichment in sorted CLL cell subpopulations: sIgMhi (red circles), sIgMint (blue squares), and sIgMlo (green triangles). Data are expressed as fraction of new cells (F %) normalised to one day’s labeling; error bars represent the standard deviation of ≥4 replicate measurements by GC-MS. Black arrows denote oral administration of deuterium labeled glucose. Yellow bars represent administration of idelalisib which was commenced 24 hours after the commencement of deuterium labeling. Time points represent days post- first deuterium labeling (day 0).

**Figure 7. Hypothetical model of leukemic cell kinetics in untreated *IGHV* mutated CLL**

Proposed model of tumor kinetics in CLL that fits the deuterium labeling data in the present study. A proliferative subset of cells transits between PB and lymphoid tissues. These cells become labeled with deuterium, change their phenotype from CXCR4loCD5hi to CXCR4intCD5int and express high levels of surface BCR. A largely distinct non-proliferating subset is also present. This subset does not label with deuterium, has a CXCR4hiCD5lo phenotype and low expression of surface BCR. These cells generally do not migrate into lymphoid tissues but, unlike normal anergic B cells, which undergo apoptosis when excluded from germinal centres, these cells express bcl-2 and so survive.

**Table 1. Baseline clinical and laboratory characteristics of patients recruited to deuterium labeling studies**

**Table 2. Comparison of models for CLL subpopulation kinetics**

Deuterium labeling data in CLL subpopulations (defined by CXCR4/CD5 expression, sIgM expression and eIgM expression) were fitted to two mathematical models: Model A, a single population of leukemic cells in peripheral blood with cells transitioning between phenotypic sub-groups, and Model B, with an independent subpopulation of non-proliferating cells.

Lower AICC (I.e. more negative) indicate a better fit to the data. The ‘winning’ model is defined by a difference in the AICc of at least 1. Differences of <1 are denoted ‘-‘. The null hypothesis that model A is correct can be rejected with certainty (p=0.008; Wilcoxon signed rank test two sided).