Research paper

Single-cell analysis reveals the KIT D816V mutation in haematopoietic stem and progenitor cells in systemic mastocytosis

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1. Introduction

Haematopoietic stem cells (HSCs) in the bone marrow give rise to blood cells and mast cells. Differentiating HSCs progress through a number of intermediate progenitors with multilineage-forming capacity before commitment to the mast cell lineage [1,2]. The binding of stem cell factor (SCF) to its receptor, KIT, promotes the maturation and proliferation of mast cells [3–6]. It is therefore not surprising that mutations in the KIT gene coincide with the mast cell-driven disease systemic mastocytosis (SM) [7].

SM is a haematological neoplasm in which infiltrates of neoplastic mast cells occur in various tissues [8,9]. The majority of SM patients carry a mutation in KIT, most commonly affecting codon 816 [10]. The KIT D816V mutation makes receptor signalling constitutively active, independent of binding to its ligand SCF. The detection of KIT D816V in either bone marrow or peripheral blood samples is one of the criteria for the clinical diagnosis of SM using a standardised qPCR assay [11]. When

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The patients included in our experimental studies were classified from patients under evaluation for SM and from control subjects.

2.2. Patients and sample preparation

In accordance with the World Medical Association Declaration of Helsinki.

The study was approved by the Stockholm and Uppsala Regional Ethics committees. Oral and written informed consent was obtained from each patient and control subject. The study was conducted in accordance with the World Medical Association Declaration of Helsinki.

We use single-cell analysis to study the distribution of the common KIT mutation within the heterogeneous CD34+ progenitor landscape and in mast cells of systemic mastocytosis patients.

Isolation of progenitor subsets followed by cell culture identifies a population of progenitors in bone marrow with increased mast cell-forming potential. Furthermore, a novel marker for aberrant mast cells in systemic mastocytosis is described.

We delineated the cellular origin of the KIT D816V mutation in single bone marrow cells in SM by combining FACS index sorting of CD34+ HSPCs with a multiplex qPCR assay. The data revealed that the D816V mutation in KIT arises in early HSPCs. The mutation burden is low but variable in multipotent and lineage-restricted progenitor populations and increases in mature mast cells. Furthermore, the present study provides more insight into haematopoiesis in SM subjects and identifies high CD45RA expression in aberrant mast cells.

2. Materials and methods

2.1. Ethical considerations

The study was approved by the Stockholm and Uppsala Regional Ethics committees. Oral and written informed consent was obtained from each patient and control subject. The study was conducted in accordance with the World Medical Association Declaration of Helsinki.

2.2. Patients and sample preparation

Bone marrow and occasional peripheral blood samples were collected from patients under evaluation for SM and from control subjects.

The patients included in our experimental studies were classified according to the World Health Organization criteria [20] as having indolent SM (ISM, n = 19), aggressive SM (ASM, n = 3), SM with associated haematological neoplasm (SM-AHN, n = 3), or other malignancies, including myelodysplastic syndrome or myeloproliferative neoplasm (MDS or MPN, n = 2). Supplementary Table 1 shows the patient characteristics. For analytic purposes, ASM and SM-AHN patients were combined in an advanced SM group (AdvSM). The controls (Ctrl) included patients with cutaneous mastocytosis (CM, n = 3) and subjects without SM (n = 6). All samples were processed on the day of collection. Red blood cells were lysed with PharmLyse buffer (BD Biosciences, Franklin Lakes, NJ) and washed with PBS containing 2% foetal calf serum (FCS, Thermo Fisher Scientific, Waltham, MA).

2.3. Cell lines

Two human mast cell lines, HMC-1.2 (RRID: CVCL_H205) and ROSA-KIT WT (RRID: CVCL_5C49), with and without the KIT D816V mutation, respectively, were cultured as previously described [21,22].

2.4. Flow cytometry and cell sorting

Bone marrow cells were stained with fluorophore-conjugated monoclonal antibodies in Brilliant Stain Buffer (BD Biosciences, Franklin Lakes, NJ). The antibodies used for staining included CD10 (clone H10a, RRID: AB_2738247), CD14 (clone M5E2, RRID: AB_393884), CD34 (clone 581, RRID: AB_2687922), CD38 (clone Hb7, RRID: AB_2313578), CD45RA (clone HI100, RRID: AB_1727497), CD90 (clone 5E10, RRID: AB_10714644), CD117 (clone 104D2, BD Biosciences cat# 33233), CD123 (clone 6H6, RRID: AB_2562068), CD133 (clone AC133, RRID: AB_2660880), and FcRRIg (clone CRA-1, RRID: AB_1227655) (all from BD Biosciences, Biolegend, San Diego, CA, or Miltenyi Biotec, Bergisch Gladbach, Germany). The live/dead cell marker DAPI (BD Bioscience) was used for the analysis of cultured cells. Cell sorting and analysis were performed using the FACSAria III and LSRFortessa systems (BD Biosciences). For cell culture experiments, cells were sorted into sterile-filtered PBS with 2% FCS following a two-step sorting protocol, using the yield followed by the purity precision mode. The analysis of 20–100 sorted cells verified the sorting purity. Single CD34+ cells were sorted into 2 μL of lysis buffer containing 20 μg/mL PCR-grade protease K (Thermo Fisher Scientific) in EB buffer (Qiagen, Hilden, Germany). These cells were sorted either immediately from the sample or following an enrichment sort. The single-cell precision mode ensured that only one cell was deposited per well. The plates were then centrifuged and stored at −80 °C until analysis. The sorting data from single cells were saved using the index sort module of the software FACSDIVA software version 8 (BD Biosciences). Flow cytometry data analysis was performed using FlowJo software version 10 (TreeStar, Ashland, OR).

2.5. Cell culture and colony-forming assays

Colonial-forming potential was evaluated by plating 500 cells in 35 × 10 mm tissue culture dishes (Thermo Fisher Scientific) in duplicate. The cells were cultured in Methocult H4434, including SCF, IL-3, EPO, and GM-CSF (STEMCELL Technologies Inc., Vancouver, Canada). Colonies were counted after 12–13 days, distinguishing erythroid, granulocyte-monocyte, and mixed colonies following the manufacturer’s recommendations. The mast cell-forming potential was assessed via liquid culture as described previously [23]. Briefly, sorted cells were cultured in medium with recombinant human (rh) IL-3 (10 ng/mL; PeproTech, Rocky Hill, NJ) and rhIL-6 (10 ng/mL; PeproTech). In some experiments, the medium was changed on day 5, and the cells were subsequently cultured with rhIL-6 (10 ng/mL) and SCF (100 ng/mL, r-metHuSCF; Swedish Orphan Biovitrum, Stockholm, Sweden), as indicated in the figure legends. Enzyme cytochemical staining of tryptase-like activity was used to assess tryptase expression [24,25]. Cell morphology was evaluated by May-Grünwald Giemsa staining (MGG; Sigma-Aldrich, St. Louis, MO).
2.6. Single-cell mutation analysis

Before the addition of lysis buffer, 96-well plates (Bio-Rad Laboratories, Hercules, CA) were UV-treated for 30 min using an UVT-B-AR UV Cabinet (Grant Instruments, Cambridge, UK). Tubes and pipettes were treated with the DNA AWAY Surface Decontaminant (Thermo Fisher Scientific). After sorting, the cells were lysed for 45 min at 55 °C, followed by a 10-min heat-inactivating step at 95 °C. Multiplex qPCR analysis was performed via a Taqmam gene expression assay (Thermo Fisher Scientific) in lysed single cells using 96-well CFX Manager (Bio-Rad Laboratories). A mutation assay was designed based on a previously published method [11] using Primer3 (v.0.4.0) which specifically detected KIT c.2447A>T (D816V) using the forward primer 5′-AGAGACTTGGCAGCCAGAAA-3′; a reverse primer matching the mutation at the 3′ position, 5′-TTAACCACTATACTTTCTCCTTGTA-3′; and the probe 5′-FAM-TCTTCTTCTCTATTGTTAGTGTG-BHQ1-3′. A control assay was designed to target a flanking exon and intron eight in GAPDH using the forward primer 5′-CTGACTTCAACAGCGACACC-3′ and the reverse primer 5′-AGAGTTGTGTGGAGGCCCTTTC-3′; and the probe 5′-HEX-TCAAGCTTCTCTCTGGTATA TGTG-BHQ1-3′ (all primers came from TAG Copenhagen, Denmark). The control and mutation assays were tested and optimised using the cell lines ROSAKIT WT and HMC-1.2. The primer concentrations, optimised for single cells, were as follows: control assay 150 nM forward/reverse primers, 100 nM probe; mutation assay 100 nM forward/reverse primers, 67 nM probe. Each plate, containing 86 forward/reverse primers, 100 nM probe; mutation assay 100 nM forward/reverse primers, 67 nM probe. Each plate, containing 86

2.7. Statistical analysis

Statistical analysis was performed using Prism software (Version 6.0 h, GraphPad Software, La Jolla, CA). The unpaired two-tailed Student’s t-test was employed when comparing two groups, and the unpaired one-way ANOVA with Tukey’s multiple comparison test was employed when comparing more than two groups, unless otherwise indicated. Statistical analysis of log-transformed data was performed for some experiments as indicated in the figure legends. Differences were considered significant when P < .05.

3. Results

3.1. Systemic mastocytosis patients have a normal CD34+ bone marrow stem and progenitor cell composition

To assess the composition of the haematopoietic cells in the bone marrow of SM patients, we designed a multicolour flow cytometry panel that identifies ten distinct HSPC subtypes and mature mast cells (Table 1; Fig. 1a-b). Colony formation assays confirmed the validity of the granulocyte-monocyte progenitor (GMP), common myeloid progenitor (CMP), and megakaryocyte-erythroid progenitor (MEP) gating strategies in SM subjects (Supplementary Fig. 1a-c). The flow cytometry data provided a specific HSPC profile for 34 bone marrow samples: 9 control, 19 ISM and 6 AdvSM (Fig. 1c; Supplementary Table 1). The percentages of HSPCs in SM samples did not significantly differ from those in control samples (Fig. 1d and Supplementary Fig. 2a). In addition, the ISM and AdvSM samples exhibited percentages of total CD34+ cells similar to the control samples (Supplementary Fig. 2b). Thus, the overall CD34+ haematopoiesis profile did not differ between the control and SM subjects.

3.2. The bone marrow CMPFCRI+ fraction presents an enhanced mast cell-forming potential

Mast cell progenitors (MCPs) circulate in the blood as Lin− CD34+ CD117+ FcεR1+ cells [33]. These cells express CD123 and lack CD45RA [23,33], overlaying the CMPFCRI+ gate in bone marrow (Supplementary Fig. 3a) that exhibit a distinct expression profile compared with mature mast cells (Supplementary Fig. 3b). We therefore analysed the mast cell-forming potential of the bone marrow CMPFCRI+ fraction, by culturing the cells with IL-3 and IL-6. Flow cytometry analysis revealed that the CMPFCRI+ cultures gave rise to a higher frequency of CD117hi FcεR1+ mast cells than the CMPFCRI− cells (Fig. 2a-c), despite overall lower CD117 expression (Supplementary Fig. 4a). May-Grünwald Giemsa and enzymatic tryptase staining showed that the CMPFCRI+ progeny generated a high frequency of granulated tryptase-positive mast cells when cultured with SCF (Fig. 2d-e and Supplementary Fig. 5). Cultured GMPs did not give rise to any cells with a mast cell phenotype (Fig. 2a-e and Supplementary Fig. 5).

The emerging model of haematopoiesis proposes that MEPs, basophils, and eosinophils develop separately from GMPs following an asymmetric cell division at the MPP stage. This process can be traced by the segregation of CD133, which is expressed in GMPs but not in progenitors with erythromyeloid output [34]. Indeed, we found that CMPFCRI+ cells exhibited only low levels of CD133 staining, similar to erythromyeloid cells (Supplementary Fig. 4b).

3.3. CD45RA is expressed on aberrant mast cells in systemic mastocytosis

The multicolour flow cytometry panel used for the gating of HSPCs was also employed to identify mast cells in bone marrow (Fig. 1b).

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**Table 1** Immunophenotypes of haematopoietic stem and progenitor cells.

| Short | Cell type | Gating strategy | References |
|-------|-----------|-----------------|------------|
| HSC   | Haematopoietic stem cell | CD14− CD34− CD38− CD45RA− CD10− CD90+ | [26,27] |
| MPP   | Multipotent progenitor | CD14− CD34+ CD38− CD45RA− CD10− | [26,27] |
| CMPFCRI− | Common myeloid progenitor FcεR1− | CD14− CD34+ CD38− CD45RA− CD10− CD123− FcεR1− | [28] |
| CMPFCRI+ | Common myeloid progenitor FcεR1+ | CD14− CD34+ CD38− CD45RA− CD10− CD123+ FcεR1+ CD117+ | [28] |
| MEP   | Megakaryocyte-erythroid progenitor | CD14− CD34+ CD38− CD45RA− CD10− CD123− CD123+ FcεR1− | [28] |
| LMPP  | Lymphoid-primed multipotent progenitor | CD14− CD34+ CD38− CD45RA− CD10− | [28] |
| GMP   | Granulocyte-monocyte progenitor | CD14− CD34+ CD38− CD45RA− CD10− CD123+ | [28] |
| CD19  | Common dendritic progenitor | CD14− CD34− CD38− CD45RA− CD10− CD123+ | [30,31] |
| MLP   | Multilymphoid progenitor | CD14− CD34− CD38− CD45RA− CD10+ | [32] |
| BNRPC | B- and NK cell progenitor | CD14− CD34− CD38− CD45RA− CD10+ | [32] |
| MC    | Mast cell | CD14− CD34− CD117+ FcεR1+ | [32] |

* Refers to the original CMP population without the inclusion of the FcεR1 marker.
May-Grünwald Giemsa and tryptase staining confirmed that CD14− CD34− CD117hi FcεRIhi cells constitute mast cells (Fig. 3a). As expected, SM samples presented higher percentages of mast cells than controls (Fig. 3b). Notably, mast cells from almost all SM samples exhibited aberrant CD45RA expression, as verified with fluorescence minus one control and internal control staining of CD34+ cells (Fig. 3c-d).

Fig. 1. Patients with systemic mastocytosis exhibit a normal composition of CD34+ haematopoietic stem and progenitor cells. (a) A landscape model of the haematopoesis. (b) Gating strategy for bone marrow HSPCs and mature mast cells, showing representative plots with population frequencies as percentages. (c) The haematopoietic progenitor profile of 34 bone marrow samples. Cutaneous mastocytosis samples are indicated in italics. (d) SM samples were compared with control samples without SM, including CM (open circles), and the mean expression ± SEM of each HSPC is presented as a percentage of the total CD34+ cells. The unpaired t-test was used in panel d to compare the control and SM groups for each HSPC subset.
cutaneous (non-systemic) mastocytosis samples and MDS/MPN samples expressed low levels of CD45RA on bone marrow mast cells (Fig. 3c-d). Taken together, SM bone marrow samples contain more mast cells than control samples and these mast cells express high levels of the aberrant marker CD45RA.

3.4. Single-cell index sorting reveals the KIT D816V mutation in early HSPCs in systemic mastocytosis patients

The cellular origin of the KIT D816V mutation in SM has not been established, and the relative abundance of this mutation in different CD34⁺ HSPC populations and mast cells has not been determined. We therefore developed a single-cell method to trace the KIT D816V mutation using two cell lines: HMC-1.2 cells carrying the mutation and ROSA cells carrying wild-type KIT. The cells were labelled with two different fluorochrome-labelled antibodies, mixed, and randomly FACS sorted with the index sort function. A multiplex qPCR assay was used to detect an irrelevant gene, for quality control purposes, and the KIT D816V mutation (Supplementary Fig. 6a-b). Among all single cells, >96% passed the quality control procedure (Supplementary Fig. 6c). Linking the results with the index data revealed that the mutation assay was negative for the wildtype ROSA cells. The assay detected the mutation in 98.1% of

Fig. 2. The CMP<sup>FceRI⁺</sup> fraction in the bone marrow has mast cell potential. (a-c) Flow cytometry analysis of CMP<sup>FceRI⁺</sup>, CMP<sup>FceRI⁻</sup> and GMP cells cultured with rhIL-3 and rhIL-6 up to 21 days. (a) Flow cytometry plots of representative cultures, showing the percentages of CD117⁻, CD117⁺, and CD117<sup>hi</sup> cells in each plot. (b) Percentage of CD117<sup>hi</sup> FceRI⁺ cells at days 5–6. (c) Two of the samples in panel b were followed for up to 21 days under culture with rhIL-3 and rhIL-6. The percentage of CD117<sup>hi</sup> cells was analysed on days 5–6, 13–14, and 19–21. (d) Representative images of CMP<sup>FceRI⁺</sup>, CMP<sup>FceRI⁻</sup> and GMP cells cultured with rhIL-3 and rhIL-6 for 5 days, followed by rhIL-6 and SCF until day 13–14, showing tryptase staining (red) and granules by May-Grünwald Giemsa (MGG) staining. (e) Quantification of tryptase-positive cells at days 13–14 and 17–21. CMP<sup>FceRI⁺</sup>, CMP<sup>FceRI⁻</sup> cells were analysed from Ctrl29, SM28, SM32, SM33, and MDS30. GMPs were analysed from SM32 and SM33. The bars and lines in panel b, c and d represent the means ± SEM. One-way ANOVA with Tukey’s multiple comparison was used in panel b, **P < .01, ***P < .001. Images were captured using an Olympus XC10 camera (Olympus, Tokyo, Japan). The image width corresponds to 29 μm.
the mutation is mainly restricted to mast cells [13].

It is of particular interest to study the mutation burden in progenitors with mast cell-forming potential. Detection of the Kit mutation in bulk-sorted CD34⁺ mast cells (Fig. 4b). Index data analysis of individual mast cells demonstrated that CD45RA expression was associated with presence of the Kit mutation (Fig. 4c). The percentage of mutated mast cells was correlated with the total percentage of mast cells in the sample and with the serum tryptase levels (Fig. 5a-b; Supplementary Table 3). The mutation rate in CD34⁺ HSPCs was <3% in all samples (Fig. 4d), except for one AdvSM sample with a high mutation burden of 37.0% (Supplementary Table 4). Nevertheless, the percentage of mutated CD34⁺ HSPCs was correlated with serum tryptase levels (Fig. 5c; Supplementary Table 3).

Coupling the index data with the mutation analysis of 10 bone marrow samples revealed that the Kit D816V mutation appeared throughout the haematopoietic landscape (Fig. 4e; Supplementary Table 2 and 4). Specifically, different HSPCs presented similar mutation rates in all samples when the median percentage of mutation was compared between different HSPCs (Fig. 4f). Overall, the method of index sorting provided a representative set of single cells from the whole-bone marrow aspirates (Supplementary Fig. 7a). However, the CMPFc⁺ population with an enriched mast cell potential was rare. To determine the mutation frequency in the CMPFc⁺ fraction, we selectively sorted 86 and 278 single CMPFc⁺ cells from 2 SM samples. One patient exhibited <2% mutated CMPFc⁺, HSCs and GMPs, whereas the fraction of mutated mast cells was 98.9% (Fig. 4g; Supplementary Table 4). The second patient presented 95.7% mutated CMPFc⁺ and 94.6% mutated mast cells (Fig. 4h; Supplementary Table 4). Peripheral blood MCPs (CD14⁺ CD34⁺ CD117⁺ FcεRI⁺) showed mutation rates comparable to that of bone marrow CMPFc⁺ in one patient (Supplementary Fig. 7b), but the mutation could not be detected in the MCPs from 4 other SM samples, likely due to the low mutation frequency and the small number of cells analysed due to the rarity of the cells (14–23 cells analysed per sample). Taken together, we detected the Kit D816V mutation throughout the haematopoietic landscape, starting with the HSCs.

4. Discussion

The cellular origin of the aberrant clone in SM is still under debate. SM patients exhibit an increased frequency of mast cells in bone marrow, and here we show that almost all these mast cells carry the Kit D816V mutation. Despite normal HSPC frequencies in bone marrow, the Kit D816V mutation can be traced all the way back to the CD34⁺ CD38⁻ CD45RA⁻ CD10⁻ HSC fraction.

The Kit D816V mutation causes receptor autoactivation, which signals survival, proliferation, and differentiation, likely explaining the competitive growth advantage of the aberrant mast cell clone and the high frequency of mutated mast cells. However, the frequency of mutated HSPCs was low in most patients. This observation is consistent with the overall normal HSPC composition of SM patients. There are two main potential explanations for the low D816V mutation burden observed in HSPCs relative to mast cells: HSPCs rapidly differentiate into mast cells when the mutation occurs, or mutated HSPCs do not present a strong growth advantage over non-mutated HSPCs. In fact, we have previously shown that Kit signalling is dispensable for mast cell progenitor development in vitro and in vivo, and that other stimuli can induce cell proliferation [23]. However, the aberrant progenitor clone may in some cases proliferate to reach a high frequency, as demonstrated in two patients with AdvSM who presented >20% mutated cells in the HSC fraction. Advanced forms of SM have previously been associated with detection of the Kit mutation in bulk-sorted CD34⁺ progenitors and various mature cell lineages, whereas in indolent disease, the mutation is mainly restricted to mast cells [13].

It is of particular interest to study the mutation burden in progenitors with mast cell-forming potential. Detection of the Kit mutation in a particular progenitor is not necessarily indicative of mast cell-forming potential. Therefore, cell culture experiments were performed to investigate the fate of putative MCPs in bone marrow. We have previously reported that CD34⁺ CD117⁺FcεRI⁺ cells in peripheral blood are committed or nearly committed MCPs [33]. Considering the
CD34+ CD117+ FcεRI+ phenotype of CMPsFcεRI+ cells, these cells might be expected to present a similar mast cell-forming capacity as to blood MCPs. In the current study, we showed that CMPsFcεRI+ gave rise to a higher frequency of mast cells than CMPsFcεRI- and GMPs. However, the mast cell frequency in cultured CMPsFcεRI+ was b10% and a substantial fraction were found to be CD117−/lo FcεRI+ cells after culture, consistent with a basophil-like phenotype. Thus, CMPsFcεRI+ exhibit a considerable mast cell-forming capacity but also include progenitors with non-mast cell output. Basophils mature in bone marrow, whereas mast cells mature in peripheral tissues, which likely explains why bone marrow CMPsFcεRI+ form CD117− FcεRI+ basophil-like cells, whereas CD34+ CD117int/hi FcεRI+ blood progenitors do not. The high frequency of CD117− FcεRI+ basophil-like cells that develop from CMPsFcεRI+ is also in agreement with the reduced CD117 expression of CMPsFcεRI+. CD117 is upregulated upon mast cell maturation, and blood mast cell progenitors express intermediate to high levels of CD117. CMPsFcεRI+ express CD117 at low levels overall, even though there are individual CMPsFcεRI+ cells that express high CD117 levels, which might correspond to the cells with mast cell-forming capacity. Taken together, the CMPFcεRI+ cells’ capacity to form not only mast cells, but also CD117− FcεRI+ basophil-like cells, likely explains the low KIT mutation prevalence in this progenitor population.

In SM, mast cells are known to aberrantly express multiple surface proteins, among CD2 and CD25 are currently used for diagnosis [35]. These markers are expressed independently of the SM subtype. Other markers, such as CD30 and CD52, are associated with AdvSM and mast cell leukemia [36,37]. In the present study, we demonstrate that CD45RA is a novel marker for aberrant mast cells in SM. Remarkably,
bone marrow mast cells from three patients with cutaneous mastocytosis expressed low CD45RA levels, suggesting that CD45RA is a marker for systemic mast cell disease. The increase in the CD45RA isoform of CD45 likely explains the high levels of CD45 observed in systemic mastocytosis patients [38–41]. Taken together, our results indicate that high CD45RA expression should be considered a novel marker for diagnosis of SM.

CD45RA is present on aberrant mast cells in SM, but at what stage during mast cell differentiation is the marker upregulated? CMPsREF, which lack CD45RA, present mast cell-forming potential in SM patients. In contrast, we show that cells in the classic CD45RA+ GMP gate lack mast cell potential in SM patients. An alternative explanation is that the mast cell progenitors in SM fall into the immature CD38− gate. In fact, examples of rare mutated CD117int/hi FcεRint/hi cells in the CD45RA− LMPP gate were seen (Supplementary Fig. 7c). It is also tempting to speculate that CD45RA is upregulated once aberrant mast cells are mature.

The CD34+ HSPC population is highly heterogeneous and a single-cell resolution is therefore necessary to delineate the cellular distribution of the mutation. Here, we used single-cell index sorting to track cellular identity when performing mutation analysis. Notably, the mutation was detected throughout differentiation, from HSCs to the MEP, GMP, and LMPP fractions, depending on the patient. The detection of the mutation in early HSPCs explains previous findings of the mutation being present in bulk-sorted mature cell lineages of some patients. Multiplex TaqMan qPCR assays detected the KIT mutation in index-sorted progenitors in the present investigation. The qPCR method was originally developed for sensitive mutation detection in DNA extracted from bone marrow or blood cells. GAPDH primers and probes were added to the assay to confirm the presence or absence of a sorted cell. This method allowed the detection of mutated cells with close to 100% efficiency, with no observed false positives. The single-cell index sorting and DNA mutation analysis approach can similarly be applied to other mutations and haematological diseases to reveal the origin of aberrant clones. One limitation of the method is the number of cells that can be analysed. Consequently, stochastic processes can influence the determination of low mutation rates, meaning that mutated cells in rare subpopulations might go undetected if the mutation rate is low. Similarly, determined mutation rates in populations with few mutated cells might be overestimating the actual frequency of mutated events. However, the conclusion that the KIT D816V can be found throughout the HSPC landscape is still valid.

More than 10,000 HSPCs were analysed in the present investigation, and single-cell index sorting coupled with qPCR allowed unbiased and prospective analysis of mutated HSPC populations in SM. Whole-genome amplification and nested PCR methods for amplifying and sequencing DNA mutations from single cells are associated with the phenomenon known as allelic dropout, which is the amplification of only a single allele, leading to an overestimation of the frequency of non-mutated cells. Mutational profiling of single colonies, as performed by Jawhar et al. [18] avoids this problem [18]. However, such analysis can be performed only on progenitors that form sufficiently large colonies. Mast cell progenitors present a poor proliferative capacity, and this technique was therefore not applied in the present investigation.

In summary, we have explored the composition and mutation profile of CD34+ HSPCs from SM patients. The results show that the KIT mutation can be found in cells throughout the haematopoietic landscape with increased burden in the mast cell lineage, supporting the notion that the aberrant clone may arise in the haematopoietic stem cell compartment. Furthermore, the mutated mast cells express CD45RA, a potential novel clinical biomarker for aberrant mast cells in SM.

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Declaration of interest

The authors declare no competing interests.

Author contributions

Conceptualization and Methodology, J.G., G.N. and J.S.D.; Investigation, J.G., M.E. and E.H.; Formal analysis, J.G.; Visualization and Validation, J.G. and J.S.D.; Resources, J.S.U., M.K., R.A., S.S., M.M. and M.A.; Writing – original draft, J.G. and J.S.D.; Writing – review & editing, all authors; Funding acquisition and Supervision, J.U., G.N. and J.S.D.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebiom.2019.03.089.
