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**Plasmodium-specific atypical memory B cells are short-lived activated B cells**

**Abstract** A subset of atypical memory B cells accumulates in malaria and several infections, autoimmune disorders and aging in both humans and mice. It has been suggested these cells are exhausted long-lived memory B cells, and their accumulation may contribute to poor acquisition of long-lasting immunity to certain chronic infections, such as malaria and HIV. Here, we generated an immunoglobulin heavy chain knock-in mouse with a BCR that recognizes MSP1 of the rodent malaria parasite, *Plasmodium chabaudi*. In combination with a mosquito-initiated *P. chabaudi* infection, we show that *Plasmodium*-specific atypical memory B cells are short-lived and disappear upon natural resolution of chronic infection. These cells show features of activation, proliferation, DNA replication, and plasmablasts. Our data demonstrate that *Plasmodium*-specific atypical memory B cells are not a subset of long-lived memory B cells, but rather short-lived activated cells, and part of a physiologic ongoing B-cell response.

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**Introduction**

Atypical memory B cells (AMB) are an unusual B-cell subset detected in both mouse models and humans in the context of certain infections and autoimmune disorders, including HIV, HCV, tuberculosis, malaria, rheumatoid arthritis and systemic lupus erythematosus, and accumulated with age (Knox et al., 2017b; Naradikian et al., 2016a; Portugal et al., 2017; Rubtsov et al., 2017). In the context of infections, AMB were first described in HIV-viremic subjects, and termed tissue-like memory B cells, due to their similarity to an FCRL4-expressing memory B-cell subset found in human tonsillar tissues (Ehrhardt et al., 2005; Moir et al., 2008). In addition to FCRL4, these cells express relatively high levels of other potentially inhibitory receptors including CD22, CD85j, CD85k, LAIR-1, CD72, and PD-1, and show a profile of trafficking receptors including expression of CD11b, CD11c and CXCR3, consistent with migration to inflamed tissues. They are antigen-experienced class-switched B cells, which lack the expression of CD21 and the hallmark human memory B-cell marker CD27. Further studies demonstrated the expression of the transcription factor T-bet and the cytokine IFNγ by these cells, also characteristic of Th1 CD4^+^ T cells (Knox et al., 2017b; Obeng-Adjei et al., 2017; Portugal et al., 2017). Due to their poor functional capacity upon in vitro re-stimulation with BCR ligands, AMB were characterized as dysfunctional B cells, and increased frequencies of these cells was proposed to be a consequence of B-cell exhaustion driven by chronic inflammation and stimulation, drawing parallels with T-cell exhaustion during chronic viral infections.
(Moir et al., 2008; Portugal et al., 2015; Sullivan et al., 2015). It has been hypothesized that expansion of AMB might contribute to the mechanisms driving autoimmune disorders and deficiencies in acquisition of immunity to chronic infections. However, due to lack of good tools and animal models to analyze antigen-specific atypical B cells in greater depth, many of these concepts remain speculative.

Several studies suggest that AMB might contribute to poor acquisition of long-term immunity to Plasmodium infection (Illingworth et al., 2013; Portugal et al., 2015; Sullivan et al., 2015; Sullivan et al., 2016; Weiss et al., 2011; Weiss et al., 2009; Weiss et al., 2010). Indeed, some studies demonstrated that in the absence of constant re-exposure, Plasmodium-specific serum antibody levels rapidly wane, and full protection from clinical symptoms is lost, suggesting that B-cell memory is functionally impaired (Portugal et al., 2013). However, others have reported long-lasting maintenance of Plasmodium-specific antibodies and/or memory B cells in settings of differing malaria endemicity, and similar responses are also observed in mouse malaria models (Dorfman et al., 2005; Ndungu et al., 2009; Ndungu et al., 2013; Ndungu et al., 2012; Wipasa et al., 2010). Moreover, it has been shown that BCRs cloned from P. falciparum-specific AMB from malaria-exposed adults encode P. falciparum-specific IgG antibodies, which could contribute to P. falciparum-specific IgG antibodies in serum (Muellenbeck et al., 2013). These authors proposed that P. falciparum-specific AMB do not prevent, but rather contribute to the control of Plasmodium infection. These apparently contradictory results may reflect the fact that some studies were performed on the general peripheral blood B-cell pool and others focused on Plasmodium-specific B cells. In determining a role for these cells in a chronic infection it would be important to follow antigen-specific responses and to distinguish these from non-specific polyclonal B cell activation.

The study of the development of AMB is challenging and requires suitable mouse models, which allow for identification and isolation of antigen-specific B cells that exist often at very low frequency. Here, we generated a knock-in transgenic mouse with a high frequency of B cells specific to the 21 kDa C-terminal fragment of Plasmodium chabaudi Merozoite Surface Protein 1 (MSP121), to investigate memory B cells generated following mosquito-transmission of the rodent malaria, P. chabaudi. We identified a CD11b+CD11c+FCRL5hi subset of MSP121-specific B cells during the chronic infection with phenotypical and transcriptional features strikingly similar to those of human AMB. These AMB disappeared as the infection progressed, leaving a CD11b−CD11c−FCRL5hi MSP121-specific B-cell compartment with characteristics of long-lived classical memory B cells (Bmem) after the resolution of the infection. These short-lived MSP121-specific AMB were also generated in response to immunization, suggesting they may be a normal but transient component of a B-cell response to antigen. In this chronic P. chabaudi infection, it appears that AMB require ongoing antigenic stimulation driven by the sub-patent infection to persist, and do not represent a true long-lived ‘memory’ B cell subset. Moreover, we show that generation of Plasmodium-specific AMB does not prevent the generation of Plasmodium-specific Bmem, and does not prevent resolution of the infection.

Results

Generation of an immunoglobulin heavy chain knock-in transgenic mouse model to study Plasmodium-specific B cell responses

To study Plasmodium-specific B cell responses in a rodent malaria model, we generated an IgHNMIMP23/+ mouse strain on the C57BL/6J background (Materials and methods and Figure 1—figure supplement 1).

The IgHNMIMP23/+ mice were healthy, with no unusual behavioral or physical characteristics. There were no alterations in total cellularity, pro-B, pre-B, immature B, mature B, total B220+CD19+ B cells, and plasma cells in the bone marrow (Figure 1A–B), and no alterations in number of T1, T2, T3, follicular, marginal zone, germinal center B cells, plasmablasts, plasma cells, and total cellularity in the spleen of IgHNMIMP23/+ mice (Figure 1C–D) (Sen et al., 1990; Young et al., 1994). Importantly, The IgHNMIMP23/+ mice had a greatly increased frequency of B cells specific for MSP121 (approximately 60% of the total B-cell compartment), as demonstrated by flow cytometry analysis of splenocytes with a MSP121 fluorescent probe consisting of biotinylated recombinant MSP121 loaded on streptavidin-PE (Figure 1E–F). Thus, in this model, a recombinant light chain is not required to bring about
Figure 1. Analysis of total bone marrow and splenic B-cell populations in Igh^{NIMP23/+} and Igh^{+/+} littermates. (A) Flow cytometry gating strategy to identify different B-cell populations in bone marrow of Igh^{NIMP23/+} mice. Arrows indicate flow of analysis. The same strategy was used for Igh^{+/+} littermates. (B) Percentages and numbers of different B-cell populations in bone marrow of Igh^{NIMP23/+} and Igh^{+/+} littermates as defined in (A). (C) Flow cytometry gating strategy to identify different B-cell populations in spleen of Igh^{NIMP23/+} mice. (D) Percentages and numbers of different B-cell populations in spleen of Igh^{NIMP23/+} and Igh^{+/+} littermates as defined in (A). (E) Gated on B220^+CD19^+ B cells (spleen) (F) Percentages of specific B-cells.
specificity. This suggest that most endogenous light chains will pair with the NIMP23 heavy chain to generate a BCR with detectable binding to MSP121.

Increase in Plasmodium-specific B cells after mosquito transmission of P. chabaudi
To investigate B cells in P. chabaudi infections, which last several weeks, and to avoid potential problems with activation arising from very high frequencies of MSP121-specific B cells, we reduced the precursor frequency of MSP121-specific B cells to match the natural level expected for antigen-specific B cells more closely, yet still readily detectable by flow cytometry. We generated mixed bone marrow (BM) chimeras by adoptively transferring a mixture of 10% bone marrow from either IghNIMP23+/+ or Igh+/+ mice (CD45.2+) together with 90% bone marrow from C57BL/6.SJL-Ptprca mice (CD45.1+) into sub-lethally irradiated Rag2−/−.C57BL/6.SJL-Ptprc−/− mice (CD45.1+) to generate NIMP23→Rag2−/− and WT→Rag2−/− bone marrow chimeric mice respectively (Figure 1—figure supplement 2A–B). In both types of chimeras, 2–3% of the B cells were CD45.2+ and in NIMP23→Rag2−/− mice approximately 1–2% of the B cells were MSP121-specific (Figure 1—figure supplement 2C–E). No MSP121-specific B cells were detected in the control WT→Rag2−/− chimeras (Figure 1—figure supplement 2D).

Infection of C57BL/6J wt mice with P. chabaudi by mosquito bite gives rise to a short (48 hr) pre-erythrocytic infection, followed by an acute blood parasitemia peaking approximately 10d post-transmission. Thereafter, the infection is rapidly controlled, reaching very low parasitemias by 15d post-transmission, with a subsequent prolonged (~90 d), but low-level chronic infection before parasite elimination (Brugat et al., 2017; Spence et al., 2013). NIMP23→Rag2−/− mice infected with P. chabaudi by mosquito bite, showed a similar course of parasitemia to that of control WT→Rag2−/− mice (Figure 1—figure supplement 2F), and C57BL/6J wt mice (Brugat et al., 2017; Spence et al., 2013; Spence et al., 2012). Importantly, the MSP121-specific IghNIMP23+/+ B cells (CD45.2+ MSP121+) in NIMP23→Rag2−/− chimeras showed a robust response to the infection, as demonstrated by a dramatic increase in the proportions and numbers of GL-7+ CD38+ germinal centers (GC) and IgG2b−IgD− class-switched B cells in the spleen at 35 days post-infection (dpi) (Figure 1—figure supplement 2G–H).

Thus, we have generated a mouse model with detectable numbers of functional MSP121-specific B cells capable of responding to P. chabaudi infection.

Generation of Plasmodium-specific AMB after mosquito transmission of P. chabaudi infection
We investigated whether Plasmodium-specific AMB could be identified in mice during a blood-stage P. chabaudi infection. We selected a series of mouse homologues to human cell surface markers described on human AMB (Charles et al., 2011; Kardava et al., 2014; Kardava et al., 2011; Knox et al., 2017a; Li et al., 2016; Moir et al., 2008; Muellenbeck et al., 2013; Portugal et al., 2015; Russell Knode et al., 2017; Sullivan et al., 2015). Human AMB express CD11b, CD11c, Fc receptor-like (FCRL) 3–5, high levels of CD80, low levels of CD21, and are Ig class-switched. Mouse FCRL5 most closely resembles human FCRL3 and is the only mouse FCRL-family member which
contains both ITIM and ITAM motifs in its cytoplasmic tail (Davis, 2007; Davis et al., 2004; Won et al., 2006; Zhu et al., 2013). Therefore, our flow cytometry panel for mouse AMB included antibodies against CD11b, CD11c, FCRL5, CD21, IgD, and also CD80 and CD273 which identify mouse B cells that are antigen-experienced and potentially memory cells (Anderson et al., 2007; Tomayko et al., 2010; Zucarro-Catania et al., 2014).

We detected an increased number of cells in a distinct CD11b+CD11c+ MSP121-specific B-cell subset at 28-35dpi, in the chronic phase of P. chabaudi infection (Figure 2A–B). This subset showed several AMB characteristics, including high expression of FCRL5 and low expression of CD21 and IgD (Figure 2C–E). In addition, the CD11b+CD11c+ MSP121-specific B-cell subset was enriched with cells expressing CD80 and CD273 (Figure 2C–E).

We then explored whether this CD11b+CD11c+ MSP121-specific B cell subset was detected during the memory phase, that is after resolution of the infection. As it takes up to 90 days for a blood-stage P. chabaudi infection to be eliminated from C57BL/6J mice (Achtman et al., 2007; Spence et al., 2013), we measured these responses from 155dpi onwards. Unexpectedly, the numbers of CD11b+CD11c+ MSP121-specific B cells were not significantly higher than background level (Figure 2A–B).

These data demonstrate that a mosquito-borne infection with P. chabaudi generates Plasmodium-specific B cells resembling human AMB. However, these cells do not persist and are not detected above background level after parasite clearance.

Transcriptome analysis confirms the AMB nature of CD11b+CD11c+ MSP121-specific B cells, and reveals a plasmablast-like signature for this subset

To gain a better understanding of the identity of the CD11b+CD11c+ Plasmodium-specific B cell subset, we isolated both CD11b+CD11c- and CD11b+CD11c+ MSP121-specific B cells from spleens of P. chabaudi-infected NIMP23→Rag2−/− mice (35dpi) (Figure 3—figure supplement 1), and MSP121-specific B cells from the spleen of naive NIMP23→Rag2−/− mice (Figure 2A), by flow cytometric sorting, and performed an mRNAseq transcriptional analysis on the three populations.

We then selected a large series of key genes previously shown to be either up or downregulated on human AMB (Supplementary file 1), and explored the expression of their mouse homologues on the three different B-cell subsets we sorted at day 35pi. The transcriptome of CD11b+CD11c+ MSP121-specific B cells highly resembled that of human AMB. A series of hallmark genes upregulated in human AMB were also upregulated in CD11b+CD11c- MSP121-specific B cells, including IgG (Ighg2c and Ighg2b), Cxcr3, Tbx21 (T-bet), Lai r1 and Fcrl5 (Figure 3, genes in red boxes, and references in Supplementary file 1). In addition, the MSP121-specific CD11b+CD11c+ B-cell subset showed upregulation of Ifng, Aicda, a large array of inhibitory receptors [including Pd1, Cd72, Cd85k, Fcgr2b (CD32b), Siglece], antigen-experienced/memory markers [Cd80, Cd86, Nt5e (CD73) and high Cd38], and additional class-switched immunoglobulins (i.e. Igha, Ighg1 and Ighg3), all of which have been shown to be upregulated on human AMB (Figure 3C–G, references in Supplementary file 1). These cells also expressed Galectins (Lgals1 and Lgals3), previously implicated in B-cell anergy (Clark et al., 2007) (Figure 3H), and displayed a pro-apoptotic program (e.g. high expression of Fasl, and low expression of Bcl2) (Figure 3I). Interestingly, in agreement with data on human AMB, MSP121-specific CD11b+CD11c+ B cells showed upregulation of Mki67 (Figure 3J), indicative of proliferation, and had characteristics of plasmablasts and/or plasma cells, including upregulation of Cd138, Prdm1 (Blimp1) and Xbp1, and low expression of Cxcr5, Pax5, and Bcl6 (Figure 3B and K). However, these cells showed low expression of Ifih1 and Sip1, suggesting that they may be in a pre-plasmablast or pre-migratory plasma-cell stage (Kabashima et al., 2006; Kallies et al., 2007) (Figure 3K). Finally, and similar to human AMB, CD11b+CD11c+ MSP121-specific B cells showed low expression of Cd40, Cr2 (CD21), Ms4a1 (CD20), and Cd24a (Figure 3L).

We ran a Gene Set Enrichment Analysis (GSEA) (Subramanian et al., 2005) with a gene list ranked according to their differential expression between MSP121-specific CD11b+CD11c+ AMB sorted from infected mice and MSP121-specific B cells sorted from naive mice, using gene sets a priori obtained from Reactome (Fabregat et al., 2018). Among the gene sets yielding the top 50 significant (fdr <0.001) highest normalized enrichment score (NES) we obtained gene sets corresponding to cell cycle, DNA replication, generation/consumption of energy, regulation of apoptosis, activation of NF-κB on B cells, and downstream signaling events of the BCR
These data further corroborate the activated and proliferative nature of MSP121-specific CD11b⁺CD11c⁺ AMB.

Taken together, these data demonstrate that CD11b⁺CD11c⁺ MSP121-specific mouse AMB present during the chronic phase of *P. chabaudi* infection are very similar to human AMB described in several chronic infections. In addition, this B-cell subset shows features of activation, proliferation,
Figure 3. Transcriptome analysis of sorted splenic MSP121-specific CD11b⁺CD11c⁺ AMB. MSP121-specific CD11b⁺CD11c⁺ (AMB) and CD11b⁻CD11c⁻ B cells were flow cytometry sorted from the spleen of NIMP23→Rag2⁻/⁻ chimeric mice at 35dpi; MSP121-specific B cells were flow cytometry sorted from the spleen of naïve NIMP23→Rag2⁻/⁻, and these three B cell populations were submitted to mRNAseq analysis. The heat maps display level of expression of selected individual genes, organized in functional clusters related to (A) Fc receptor like molecules, (B) cell trafficking, (C) cytokines, (D) transcription factors, (E) inhibitory receptors, (F) antigen experience/memory, (G) immunoglobulins, (H) galectins, (I) apoptosis, (J) proliferation, (K) plasma cells/plasmablasts/germinal centers, (L) surface markers. Each column corresponds to data from an individual mouse (n = 5 35 dpi, n = 5 0 dpi).

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The following figure supplements are available for figure 3:

Figure supplement 1. Gating strategy for the sorting of splenic MSP121-specific CD11b⁺CD11c⁺ AMB and CD11b⁻CD11c⁻ B cells at 35dpi.

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Figure supplement 2. Profile of the Running ES Score and positions of GeneSet members on the rank ordered list for selected Reactome pathway gene sets.

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DNA replication and plasmablasts, resembling previous observations in human AMB (Muellenbeck et al., 2013).

**Generation of Plasmodium-specific AMB in response to immunization**

The occurrence of CD11b⁺CD11c⁺ AMB might be a consequence of aberrant B-cell activation driven exclusively by certain pathogens. Alternatively, they might be part of a normal B-cell response, which is exacerbated by the persistent nature of certain infections. To test whether CD11b⁺CD11c⁺FCRL5⁺ AMB could be generated in the absence of persistent infection, we immunized mice with MSP1₂₁. A previous report had demonstrated the presence of CD11b⁺CD11c⁺Tbet⁺ B-cells 24 hr post-immunization with R848, a TLR7/8 ligand (Rubtsova et al., 2013). Therefore, we immunized IghᴺIMP²³/+ mice with R848 together with the antigen MSP1₂₁ and looked for the appearance of MSP1₂₁-specific CD11b⁺CD11c⁺FCRL5⁺ atypical B cells. We observed substantial numbers of MSP1₂₁-specific CD11b⁺CD11c⁺ B cells in the spleens of IghᴺIMP²³/+ mice 24 hr post-immunization (Figure 4A). These cells expressed increased levels of both FCRL5 and CD80 (Figure 4B–C) and did not display GC characteristics (Figure 4D), similar to the MSP1₂₁-specific CD11b⁺CD11c⁺FCRL5⁺ atypical B cells generated following *Plasmodium* infection. The MSP1₂₁-specific CD11b⁺CD11c⁺ B cells observed

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**Figure 4.** Generation of splenic MSP1₂₁-specific CD11b⁺CD11c⁺AMB in response to immunization. (A) Flow cytometry showing differential expression of CD11b and CD11c on splenic MSP1₂₁-specific B cells from IghᴺIMP²³/+ mice before immunization (day 0) and at days 1 and 3 post-immunization with R848 and MSP1₂₁. (B) Flow cytometry showing expression of FCRL5 and CD80 on different subsets of splenic MSP1₂₁-specific B cells from IghᴺIMP²³/+ defined based on CD11b and CD11c expression at day one post-immunization and naive mice. (C) Geometric MFI of FCRL5 and CD80 expression on different subsets of splenic MSP1₂₁-specific B cells from IghᴺIMP²³/+ defined based on CD11b and CD11c expression at day one post-immunization. Two-way ANOVA vs CD11b⁻CD11c⁻ subset. ****p<0.0001. (D) Flow cytometry of CD38 vs GL-7 (GC markers) on CD11b⁺CD11c⁺ B cells from IghᴺIMP²³/+ at day one post-immunization. (E) Numbers of splenic CD11b⁺CD11c⁺ MSP1₂₁-specific B cells from IghᴺIMP²³/+ during the course of immunization. Kruskal-Wallis test compared to day 0. **p<0.01. Error bars are SEM. Data pooled from three independent experiments with 3–5 mice per group.

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after immunization appeared only transiently, as they could no longer be detected at 3 and 7d post-immunization (Figure 4A and E).

These data demonstrate that MSP121-specific CD11b+CD11c+ AMB with no functional characteristics of memory B cells can be generated independently of the infection and the presence of the pathogen, and that they are short-lived cells.

**Plasmodium-specific CD80⁺CD273⁺ Bₘₐₐₜ are generated and persist after resolution of *P. chabaudi* infection**

Identification of mouse Bₘₐₐₜ by flow cytometry originally relied on detecting B cells that had undergone Ig class-switching from IgM to IgG, and that did not express GC markers (i.e. IgG/CD38⁺GL-7⁻) (Lalor et al., 1992; Ridderstad and Tarlinton, 1998). More recently, this set of markers has been extended to include CD80, CD273 (PD-L2) and CD73, with CD273 and CD80 being the most useful to discriminate memory from naïve B cells (Tomayko et al., 2010; Zuccarino-Catania et al., 2014). In combination, these markers allow the identification of different subsets of switched as well as non-class switched (i.e. IgM/D⁺) Bₘₐₐₜ. Therefore, we used cell surface expression of CD80 and CD273 on MSP121-specific B cells to identify Bₘₐₐₜ during and after resolution of *P. chabaudi* infection.

MSP121-specific B cells from spleens of naïve NIMP23→Rag2⁻/⁻ mixed BM chimeras showed little expression of either CD80 or CD273 (Figure 5A). By contrast, CD80⁺CD273⁺, CD80⁻CD273⁻ and CD80⁻CD273⁺ MSP121-specific B cells, both class-switched (IgM/D⁺) and non-class-switched (IgM/D⁻), were readily detected above background levels at 28-35dpi in *P. chabaudi*-infected NIMP23→Rag2⁻/⁻ mixed BM chimeras (Figure 5B and D). After resolution of infection (155-170dpi), the numbers of CD80⁺CD273⁺ and CD80⁻CD273⁻ MSP121-specific B cells were either sustained or increased, while the CD80⁻CD273⁺ population decreased, compared with 28-35dpi (Figure 5A–D). All three MSP121-specific Bₘₐₐₜ subsets showed high CD38 expression (Figure 5E,F). Importantly, the MSP121-specific GC B cells detected at 28-35dpi did not express either CD80 or CD273, further distinguishing GC from memory and atypical memory P. chabaudi-specific B-cell subsets (Figure 5G). No MSP121-specific GC B cells were detected above background level after resolution of the infection (Figure 5H). Finally, and in accordance with Figure 3, approximately 70% of the MSP121-specific CD11b⁺CD11c⁺ AMB observed at days 28-35pi expressed either CD80, CD273 or a combination of both (Figure 5I).

These data show that, in contrast to the transient AMB, splenic CD80⁺CD273⁺ and CD80⁻CD273⁻ class-switched and non-class-switched MSP121-specific Bₘₐₐₜ persist after resolution of *P. chabaudi* infection.

**Plasmodium-specific Bₘₐₐₜ express high levels of FCRL5**

As discussed above, no single marker has been described so far that can identify all mouse Bₘₐₐₜ subsets. Surprisingly, we observed that after resolution of infection (155-170dpi), MSP121-specific B cells expressing different combinations of CD80 and CD273 (CD80⁺CD273⁺, CD80⁻CD273⁻ or CD80⁻CD273⁺) all expressed very high levels of FCRL5, in contrast to CD80⁻CD273⁻ MSP121-specific B cells that express no memory markers at this stage (Figure 6A). This suggests that FCRL5 might be a marker for all Bₘₐₐₜ. In order to confirm this, we used unsupervised methods to analyze our multiparameter flow cytometry data. We used PhenoGraph and t-SNE within the Cytofkit package (Materials and methods, Chen et al 2016) to analyze MSP121-specific B cells based on the expression of FCRL5, CD38, IgD, CD80 and CD273 on these cells, as determined by flow cytometry (Figure 6A and B). The analysis identified six clusters of cells with memory characteristics displaying high expression of CD38, CD80 and/or CD273, and variable expression of IgD, all of which expressed high levels of FCRL5 (Figure 6C: clusters identified with purple arrows). We then used Isomap, (Cytofkit package) to infer the relatedness between those cell subsets identified by PhenoGraph. This confirmed high similarities between the cell clusters expressing high levels of FCRL5 with the clusters expressing high levels of the memory markers CD80, CD273 and CD38 (Figure 6D).

To confirm the memory identity of MSP121-specific FCRL5⁺ B-cells detected after resolution of the infection, we isolated MSP121-specific B cells expressing either high levels of FCRL5 or not expressing FCRL5 (i.e. FCRL5⁺ and FCRL5⁻ MSP121-specific B cells) from the spleen of *P. chabaudi*-infected NIMP23→Rag2⁻/⁻ mice (155dpi) (Figure 6—figure supplement 1), and MSP121-specific B
Figure 5. Detection of MSP121-specific B cells after resolution of P. chabaudi infection. (A), (B) and (C) Flow cytometry showing gating strategy to identify splenic IgM/Dhi and IgM/Dloi CD273+ and/or CD80+ MSP121-specific B cells in NIMP23→Rag2−/− chimeric mice before infection (day 0), at 35 and 155dpi, respectively. (D) Numbers of splenic IgM/Dhi and IgM/Dloi CD273+ and/or CD80+ MSP121-specific B cells in NIMP23→Rag2−/− chimeric mice during the course of mosquito transmitted P. chabaudi infection. Two-way ANOVA vs day 0. **p<0.01; ***p<0.001; ****p<0.0001. (E) Flow cytometry showing expression of CD38 on different subsets of splenic IgM/Dhi and IgM/Dloi MSP121-specific B cells from NIMP23→Rag2−/− chimeric mice defined based on CD273 and CD80 expression at 155dpi. (F) Geometric MFI of CD38 expression on different subsets of IgM/Dhi and IgM/Dloi splenic MSP121-specific B cells from NIMP23→Rag2−/− chimeric mice expressed at day 155 post-mosquito transmitted P. chabaudi infection. Two-way ANOVA vs CD273 CD80− subset. **p<0.01; ***p<0.001; ****p<0.0001. Error bars are SEM. (G) Flow cytometry of CD273 and CD80 expression on non-GC (CD38hiGL-7lo, blue) and GC (CD38loGL-7hi, red) splenic MSP121-specific B cells from NIMP23→Rag2−/− chimeric mice at 35dpi. (H) Flow cytometry of CD38 vs GL-7 (GC markers) on splenic MSP121-specific B cells from NIMP23→Rag2−/− chimeric mice at 0 and 155dpi. (I) CD11b+CD11c+ MSP121-specific B cells overlaid on the CD80 vs CD273 plot corresponding to total MSP121-specific B cells. Data pooled from three independent experiments with 3–7 mice per group.

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cells from the spleen of naïve NIMP23→Rag2−/− mice, by flow cytometric sorting, and performed mRNAseq analysis on these three sorted cell populations (Figure 6E). As expected, the MSP121−specific FCRL5hi B cell subset showed high expression of genes encoding the hallmark memory B cell markers Cd38, Cd80, Cd86, Nt5e (CD73) and Pdcd1lg2 (CD273), when compared with either MSP121−specific FCRL5− B cells sorted at the same time or MSP121−specific B cells sorted from naïve mice (Figure 6E). Moreover, the MSP121−specific FCRL5hi B cells sorted after resolution of the infection upregulated the anti-apoptotic Bcl2 gene, which is an additional hallmark characteristic of memory B cells (Figure 6E). Importantly, FCRL5 also identified CD80+ and CD273+ MSP121−specific Bmem subsets generated following immunization with a model antigen (Figure 6—figure supplement 2).
Thus, after resolution of the infection, high expression of FCRL5 identifies *P. chabaudi*-specific B\textsubscript{mem}.

**Plasmodium-specific AMB are a distinct short-lived activated B cell subset**

After identifying and sorting MSP1\textsubscript{21}-specific AMB during chronic *P. chabaudi* infection, and MSP1\textsubscript{21}-specific B\textsubscript{mem} after resolution of the infection, we then compared the transcriptome of these two B-cell subsets. Principal component analysis (PCA) demonstrated a strikingly distinct transcriptome of MSP1\textsubscript{21}-specific AMB from that of MSP1\textsubscript{21}-specific B\textsubscript{mem}, as well as all other MSP1\textsubscript{21}-specific B-cell subsets sorted in this study (*Figure 7A*). The MSP1\textsubscript{21}-specific AMB sorted at 35dpi formed a separated cluster at the extreme right of the PCA axis of the PCA plot, which accounts for the majority of the variance (*Figure 7A*). All the other subsets [including MSP1\textsubscript{21}-specific CD11b\textsuperscript{+}CD11c\textsuperscript{−} B-cells sorted from the same mice and at the same day post-infection as the MSP1\textsubscript{21}-specific AMB (i.e. 35dpi)] clustered on the left of the PC1 axis, and showed differences mostly along the PC2 axis of the PCA plot, which accounts for only 10% of the variance (*Figure 7A*). Interestingly, MSP1\textsubscript{21}-specific CD11b\textsuperscript{−}CD11c\textsuperscript{−} and MSP1\textsubscript{21}-specific B\textsubscript{mem} clustered on opposite sides of the MSP1\textsubscript{21}-specific naïve B-cell subset along the PC2 axis (*Figure 7A*), which suggests that the MSP1\textsubscript{21}-specific B\textsubscript{mem} more closely resemble MSP1\textsubscript{21}-specific naïve B cells than MSP1\textsubscript{21}-specific CD11b\textsuperscript{−}CD11c\textsuperscript{−} B cells sorted at 35dpi.

MSP1\textsubscript{21}-specific AMB sorted during chronic *P. chabaudi* infection, and MSP1\textsubscript{21}-specific B\textsubscript{mem} sorted after resolution of the infection shared the expression of a series of mouse memory markers, including Cd80, Fcrl5, Nt5e (CD73), and Cd86 (*Figure 7B*). However, these two subsets showed differences in the expression pattern of anti- and pro-apoptotic genes (*Figure 7C*). While MSP1\textsubscript{21}-specific B\textsubscript{mem} from after infection resolution showed the highest levels of expression of the anti-apoptotic Bcl2 gene, MSP1\textsubscript{21}-specific AMB sorted during chronic *P. chabaudi* infection showed the lowest levels of expression of this hallmark anti-apoptotic gene (*Figure 7C*). In contrast to MSP1\textsubscript{21}-specific B\textsubscript{mem}, MSP1\textsubscript{21}-specific AMB expressed high levels of the pro-apoptotic genes Bad, Bax, Fas and Fasl (*Figure 7C*). Interestingly, MSP1\textsubscript{21}-specific AMB expressed very high levels of class-switched immunoglobulins, including Igha, Ighg1, Ighg2b, Ighg2c and Ighg3 (*Figure 7D*). Finally, MSP1\textsubscript{21}-specific AMB highly expressed Cd11b, Cd11c, Tbx21, Ifng and Pdcd1 (*Figure 7E*), all hallmarks of human AMB, as well as Mki67, indicative of active cell division, as previously shown in human AMB.

These data demonstrate that AMB and B\textsubscript{mem} share the expression of memory markers. However, they show striking differences in the expression of pro- and anti-apoptotic genes, immunoglobulins genes, and cell proliferation genes. The increased expression of Mki67, pro-apoptotic genes and class-switched immunoglobulins in AMB suggests that they resemble activated B cells. By contrast, B\textsubscript{mem} express much less Mki67 (similar to naïve cells) and present an anti-apoptotic gene expression pattern, consistent with being long-lived quiescent B cells.

Mouse FCRL5 has been shown to be expressed on marginal zone (MZ) and B1 B cells (*Won et al., 2006*). Therefore, it is possible that the AMB identified in this study might represent a specific subset of either MZ or B1 B cells. As discussed elsewhere (*Baumgarth, 2011*; *Garraud et al., 2012; Pillai et al., 2005; Zouali, 2011*), B1 and MZ B cells are CD1d\textsuperscript{mid/hi}, CD9\textsuperscript{+}, IgM\textsuperscript{hi} and CD23\textsuperscript{−}. B1 are further CD43\textsuperscript{+}, B220\textsuperscript{−}, and may (B1a) or may not (B1b) express CD5. MZ are further characterised by CD22\textsuperscript{hi}, CD21/CR2\textsuperscript{hi}, and the expression of the lymphosphphilid sphingosine-1 phosphate receptor S1P\textsubscript{1}, and the lineage master regulator Notch2. Indeed, MSP1\textsubscript{21}-specific CD11b\textsuperscript{−}CD11c\textsuperscript{−} AMB showed high expression of some markers associated with B1 and/or MZ, including CD5, CD9 and CD43. However, these markers have also been shown to be highly expressed by activated B2 B cells and plasma cells (*Baumgarth, 2011*). Studying all these canonical markers, we found no other similarities between B1/MZ and MSP1\textsubscript{21}-specific CD11b\textsuperscript{−}CD11c\textsuperscript{−} AMB to suggest a relationship between these subsets (*Figure 8A*). This was further corroborated by flow cytometry analysis (*Figure 8B*). Moreover, unlike MZ and B1 B cells, MSP1\textsubscript{21}-specific AMB were class-switched and expressed very high levels of IgG (*Figure 7D*). To further characterize the identity of MSP1\textsubscript{21}-specific CD11b\textsuperscript{−}CD11c\textsuperscript{−} AMB, we explored if these cells resemble GC B cells. Flow cytometry analysis demonstrated that MSP1\textsubscript{21}-specific AMB did not display a GC phenotype (i.e. CD38\textsuperscript{hi}GL-7\textsuperscript{−}) (*Figure 8C*) while the MSP1\textsubscript{21}-specific CD38\textsuperscript{lo}GL-7\textsuperscript{hi} GC B cells showed low expression of both CD11b and CD11c. Thus, MSP1\textsubscript{21}-specific AMB are not GC B cells.
Figure 7. MSP121-specific AMB are a distinct short-lived activated B cell subset. (A) Principal component analysis of RNAseq transcriptome data from splenic MSP121-specific AMB (CD11b CD11c, 35dpi), CD11b CD11c AMB (35dpi), CD11b CD11c B cells (35dpi), Bmem (FCRL5hi, 155dpi), FCRL5 B cells (155dpi) and B cells from naive mice (0dpi). The MSP121-specific AMB and Bmem are contained inside ellipses. (B) (C) (D) (E) Normalized counts corresponding to selected Figure 7 continued on next page
genes representing memory B-cell markers, anti and pro-apoptotic genes, immunoglobulins and atypical memory B-cell markers, respectively, for all five groups described in (A). Each bar represents an individual mouse. Data generated with five mice per group.

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Figure 7 continued

All together, these data further show that *P. chabaudi*-specific AMB represent a distinct subset of short-lived activated B cells.

Discussion

Similar to other chronic infections [e.g. HIV, HCV and *Mycobacterium tuberculosis* (Knox et al., 2017b; Portugal et al., 2017)], *Plasmodium* infection, the cause of malaria, leads to an increase in the frequency of AMB (originally termed tissue-like memory B cells (Moir et al., 2008)) in peripheral blood from *P. falciparum*-exposed subjects (Illingworth et al., 2013; Portugal et al., 2015; Sullivan et al., 2016; Sullivan et al., 2015; Weiss et al., 2011; Weiss et al., 2010; Weiss et al., 2009). However, mouse models to investigate *Plasmodium*-specific AMB are lacking. Here we have generated an IgH knock-in transgenic mouse strain to study the generation of *Plasmodium*-specific AMB in a *Plasmodium chabaudi* infection. We demonstrate the generation of *P. chabaudi*-specific AMB in response to blood-stage mosquito-transmitted chronic *P. chabaudi* infection, and show the short-lived nature of these cells. *P. chabaudi* infections in mice present some outstanding characteristics for the study of AMB; a chronic phase which allows investigation of the impact of constant immune activation driven by persistent subpatent parasitemia, followed by a clearance phase which allows the study of immune responses after the infection is naturally resolved. Thus, it is possible to study both exhaustion driven by chronic immune activation, and memory immune responses which remain after *P. chabaudi* elimination.

The *P. chabaudi*-specific AMB we detected during the chronic phase of infection showed strong similarities to human AMB described in chronic malaria. These included being class-switched, and expressing mouse homologues of hallmark human atypical memory B-cell genes such as *Itgax* (Cd11b), *Itgam* (Cd11c), *Cxc3*, *Fcr5*, *Tbx21* (T-bet), *Ifng*, *Cd80*, *Cd86*, *Aicda*, and a series on inhibitory receptors, including *Lair1* and *Pdcd1* (PD-1). Human AMB have been shown to share several characteristics with plasmablasts (Knox et al., 2017a; Muellenbeck et al., 2013; Obeng-Adjei et al., 2017; Sullivan et al., 2015). Interestingly, we also observed increased expression of several genes associated with plasmablasts in *P. chabaudi*-specific AMB, including *Cd138*, *Xbp1*, *Prdm1* (encoding Blimp-1) and *Mki67*, accompanied by downregulation of *Pax5* and *Bcl6*.

AMB resemble age-associated B cells (ABC) which accumulate with age as well as in autoimmunity, and were also proposed to be a subset of long-lived memory B cells (Naradikian et al., 2016a; Portugal et al., 2017; Rubtsov et al., 2017). Expression of T-bet, CD11c and CXCR3 are shared by AMB, tissue-like memory B cells, and ABC (Knox et al., 2017b; Naradikian et al., 2016a). Moreover, similar to ABC, expansion of human AMB associated with malaria is driven by IFNγ (Obeng-Adjei et al., 2017). IL-21, which is highly expressed by follicular helper T cells in response to *Plasmodium* infection (Carpio et al., 2015; Obeng-Adjei et al., 2015; Pérez-Mazliah et al., 2015), also directly promotes T-bet expression in B cells in the context of TLR engagement (Naradikian et al., 2016b). Taken together, these data strongly suggest that this is the same T-bet+ B-cell subset, which accumulates with time due to repetitive antigenic exposure. In agreement with previous data (Rubtsova et al., 2013), we show here that immunization with MSP121 and R848, a TLR7/8 ligand, promotes a robust but short-lived CD11b+CD11c+ *P. chabaudi*-specific AMB response. T-bet+ atypical B cells are critical to eradicate various murine viral infections (Barnett et al., 2016; Rubtsova et al., 2013), and a recent study showed that yellow fever and vaccinia vaccinations of humans stimulated an acute T-bet+ B-cell response and suggested that these T-bet+ B-cell population may function as an early responder during acute viral infections (Knox et al., 2017a). Thus, T-bet+ B cells, even in the context of malaria, are likely to be a normal component of the immune compartment that becomes activated and expands, most probably in response to BCR, endosomal TLR, and IFNγ or IL-21 stimulation. Moreover, a recent study shows a TLR9/IFNγ-dependent activation of autoreactive T-bet+CD11c+ atypical B cells in response to *P. yoelii* 17XNL infection in mice.
**Figure 8.** Analysis of MZ, B1 and GC B cell characteristics on MSP1$_{21}$-specific AMB. (A) MSP1$_{21}$-specific CD11b$^+$CD11c$^+$ (AMB) and CD11b$^-$CD11c$^-$ B cells were flow cytometry sorted from the spleen of NIMP23$^+$Rag2$^-/-$ chimeric mice at 35dpi; MSP1$_{21}$-specific B cells were flow cytometry sorted from the spleen of naïve NIMP23$^+$Rag2$^-/-$, and these three B cell populations were submitted to mRNAseq analysis. The heat map displays level of expression of selected individual genes known to be up or downregulated on either MZ, B1 B cells or both. Each column represents an individual mouse. (B) Flow cytometry analysis of surface markers of either MZ, B1 B cells or both, on MSP1$_{21}$-specific CD11b$^+$CD11c$^+$ (AMB) (blue) and MSP1$_{21}$-specific IgM$^+$ (red) B cells from the spleen of Igh$^NIMP23^+$/+ mice at 20dpi. (C) Flow cytometry analysis of GC markers on MSP1$_{21}$-specific CD11b$^-$CD11c$^-$ (non-AMB, left) and CD11b$^+$CD11c$^+$ (AMB, right) B cells from the spleen of Igh$^NIMP23^+$/+ mice at 20dpi. (D) Flow cytometry analysis showing the expression of CD11b and CD11c on MSP1$_{21}$-specific CD11b$^+$CD11c$^+$ (AMB, blue) compared to GC (CD38$^+$GL-7$^+$, red) B cells from the spleen of Igh$^NIMP23^+$/+ mice. Data generated with 5–6 mice per group.

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Here, we show that *P. chabaudi*-specific AMB are short-lived activated B cells. These cells were absent after resolution of the infection, and immunization with purified antigen and TLR agonists resulted in a transient, yet robust, activation of *P. chabaudi*-specific AMB which lasted no more than 48 hr. Moreover, the IgδNIMP23/+ mouse model allowed us to obtain a deep insight of the transcriptome profile of MSP121-specific AMB during natural infection, and compare it side-by-side with the transcriptome of MSP121-specific naïve B cells. This allowed us to demonstrate their heavily pro-apoptotic and activated transcription profile, further explaining their short-lived nature. *P. chabaudi*-specific AMB showed very low expression of Bcl2, and high levels of expression of several pro-apoptotic genes including Bad, Bax, Fas and Fasl. In addition, these cells expressed very high levels of class-switched immunoglobulins and genes associated with DNA replication and proliferation.

The association of *P. chabaudi*-specific AMB with ongoing infection explains several observations in human studies: reduction of HIV plasma viremia by ART resulted in a significant reduction of HIV-specific AMB without altering the frequency of HIV-specific Bmem (de Bree et al., 2017; Kardava et al., 2014); individuals living in high malaria endemicity present higher frequencies of AMB than individuals living in areas with moderate transmission (Illingworth et al., 2013; Sullivan et al., 2015); repetitive *Plasmodium* episodes result in higher frequencies of AMB (Obeng-Adjei et al., 2017); the percentage of AMB is larger in children with persistent asymptomatic *Plasmodium falciparum* parasitemia as compared with parasite-free children (Weiss et al., 2009); previously exposed subjects significantly reduce the frequency of AMB following a year of continuous absence of exposure to *Plasmodium falciparum* infection (Ayieko et al., 2013). These observations all support the view that constant immune activation rather than impaired memory function leads to the accumulation of AMB in malaria.

After resolution of infection, *P. chabaudi*-specific AMB did not persist, but instead subsets of *P. chabaudi*-specific Bmem were readily detected. These cells expressed different combinations of previously described mouse B-cell memory markers (i.e. CD80, CD273 and CD73 (Anderson et al., 2007; Tomayko et al., 2010; Zuccarino-Catania et al., 2014)). *P. chabaudi*-specific Bmem included both class-switched and non-class-switched cells, which show different responses to a secondary challenge infection (Krishnamurty et al., 2016). Independently of the combination of previously described memory markers expressed on *P. chabaudi*-specific Bmem, all of these cells displayed very high expression of FCRL5. Previous data showed most prominent expression of FCRL5 in marginal zone B cells, while much less evident in the newly-formed and follicular splenic B cell subpopulations (Davis et al., 2004; Won et al., 2006). In agreement with this, we observed expression of FCRL5 on a subset of splenic *P. chabaudi*-specific B cells obtained from naïve mice. However, the level of FCRL5 expression on *P. chabaudi*-specific Bmem detected after resolution of the infection was noticeably higher than that of naïve B cells both at the protein and mRNA levels. In contrast to MSP121-specific AMB, these CD11b+CD11c−FCRL5+ MSP121-specific Bmem showed high expression of the hallmark memory and anti-apoptotic gene Bcl2 (Bhattacharya et al., 2007). Thus, after resolution of the infection, high expression of FCRL5 acted as a universal Bmem marker. Due to its complex dual ITIM/ITAM signaling capacity (Zhu et al., 2013), it is tempting to speculate that FCRL5 might serve as an important signal in the differentiation/maintenance of Bmem.

Tracking the fate of the different MSP121-specific B-cell subsets identified in this work will allow detailing the interplay between them. A model can be proposed in which antigen-specific AMB serve as an intermediate stage of differentiation between naïve and Bmem. Alternatively, antigen-specific AMB might represent early plasmablasts or recent GC emigrates, based on class-switching and the high expression of IgG by these cells. These scenarios are not necessarily antagonist, and might even occur in parallel. Moreover, B1 B cells have been shown to class-switch and contribute to serum IgG1, IgG2a and IgA to influenza (Baumgarth et al., 2005), and IgG-producing B1a B cells have been shown to accumulate in the spleen of a mouse model of systemic lupus erythematosus (Enghard et al., 2010). Therefore, we can’t rule out the possibility that AMB might represent a particular B1 B cell subset that expands in the spleen and blood in response to *Plasmodium* infection.

Our data suggest that the expansion of AMB in malaria is not a consequence of B-cell exhaustion, but rather a physiologic stage of B-cell activation, and that these cells are sustained in high frequencies by ongoing chronic infections. Thus, *Plasmodium*-specific AMB are neither ‘memory’, nor ‘atypical’. Importantly, our data demonstrate that robust expansion of *Plasmodium*-specific AMB does not
hinder clearance of the infection, activation of germinal centers, or generation of Plasmodium-specific long-lived quiescent B\textsubscript{mem} upon resolution of the infection.

### Materials and methods

**Key resources table**

| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|------------------------|
| Genetic reagent (M. musculus)     | B6.SJL-Ptprc\textsuperscript{a} Pepc\textsuperscript{b}/BoyJ (B6.CD45.1) | The Jackson Laboratory | MGI:4819849 | Bred in the specific pathogen-free facilities of the MRC National Institute for Medical Research and The Francis Crick Institute |
| Genetic reagent (M. musculus)     | Rag2\textsuperscript{tm1Fwa} (Rag2\textsuperscript{−/−}) | The Jackson Laboratory | MGI:1858556 | Bred in the specific pathogen-free facilities of the MRC National Institute for Medical Research and The Francis Crick Institute |
| Genetic reagent (M. musculus)     | Ig\textsuperscript{H}\textsuperscript{NIMP23+/−} | This paper | – | Bred in the specific pathogen-free facilities of the MRC National Institute for Medical Research and The Francis Crick Institute |
| Strain, strain background (Plasmodium chabaudi chabaudi, strain AS) | P. chabaudi | other | – | European Malaria Reagent Repository, University of Edinburgh |
| Strain, strain background (Anopheles stephensi, strain SD500, female) | mosquitos | PMID: 23217144 | – | Bred in Jean Langhorne’s lab |
| Antibody                          | Monoclonal Rat Anti-CD11b | BD Biosciences | 563553 | (dil 1/50) |
| Antibody                          | Monoclonal Hamster Anti-Mouse CD11c | BD Biosciences | 561022 | (dil 1/50) |
| Antibody                          | Monoclonal Rat Anti-Mouse CD138 | BD Biosciences | 553714 | (dil 1/400) |
| Antibody                          | Monoclonal Rat Anti-Mouse CD19 | BD Biosciences | 565076 | (dil 1/200) |
| Antibody                          | Monoclonal Rat anti-Mouse CD19 | Biolegend | 115530 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD19 | Biolegend | 115543 | (dil 1/400) |

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| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|--------------------------|
| Antibody                          | Monoclonal Rat anti-Mouse CD1d | Biolegend | 123510 | (dil 1/100) |
| Antibody                          | Monoclonal Rat anti-mouse CD2 | Biolegend | 100112 | (dil 1/100) |
| Antibody                          | Monoclonal Rat Anti-Mouse CD21/35 | BD Biosciences | 563176 | (dil 1/100) |
| Antibody                          | Monoclonal Rat Anti-Mouse CD21/35 | BD Biosciences | 553818 | (dil 1/100) |
| Antibody                          | Monoclonal Rat anti-Mouse CD23 | eBioscience | 25–0232 | (dil 1/100) |
| Antibody                          | Monoclonal Rat anti-Mouse CD273 | BD Biosciences | 564245 | (dil 1/25) |
| Antibody                          | Armenian Hamster anti-Mouse CD3 | Biolegend | 100336 | (dil 1/100) |
| Antibody                          | Monoclonal Rat anti-Mouse CD38 | Biolegend | 102718 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD38 | eBioscience | 17–0381 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD38 | BD Biosciences | 740697 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD4 | Biolegend | 100414 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD45.1 | Biolegend | 110706 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD45.1 | Biolegend | 110728 | (dil 1/400) |
| Antibody                          | Monoclonal Mouse anti-Mouse CD45.2 | BD Biosciences | 563685 | (dil 1/50) |
| Antibody                          | Monoclonal Mouse anti-Mouse CD45.2 | Biolegend | 109814 | (dil 1/50) |
| Antibody                          | Monoclonal Mouse anti-Mouse CD45.2 | Biolegend | 109808 | (dil 1/50) |
| Antibody                          | Monoclonal Rat CD45R/B220 | BD Biosciences | 564449 | (dil 1/400) |
| Antibody                          | Monoclonal Rat anti-Mouse CD45R/B220 | Biolegend | 103224 | (dil 1/400) |

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| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|---------------------|-------------|------------------------|
| Antibody                         | Monoclonal Rat anti-Mouse CD45R/B220 | eBioscience | 25–0452 | (dil 1/400) |
| Antibody                         | Monoclonal Rat anti-Mouse CD73 | BD Biosciences | 550741 | (dil 1/100) |
| Antibody                         | Monoclonal Armenian Hamster anti-Mouse CD80 | Biolegend | 104729 | (dil 1/25) |
| Antibody                         | Monoclonal Rat anti-Mouse CD8a | Biolegend | 100734 | (dil 1/400) |
| Antibody                         | Monoclonal Rat anti-Mouse CD9 | BD Biosciences | 558749 | (dil 1/100) |
| Antibody                         | Monoclonal Rat anti-Mouse CD93 (AA4.1) | eBioscience | 17–5892 | (dil 1/100) |
| Antibody                         | FCRL5 | PMID: 17082595 | -- | Produced in Randall Davis’ lab (dil 1/400) |
| Antibody                         | Polyclonal Sheep anti-Mouse FCRL5 | R and D Systems | FAB6756G | (dil 1/50) |
| Antibody                         | Monoclonal Rat Anti-Mouse T- and B-Cell Activation Antigen GL7 | BD Biosciences | 562080 | (dil 1/100) |
| Antibody                         | Monoclonal Rat Anti-Mouse IgD | Biolegend | 405725 | (dil 1/400) |
| Antibody                         | Monoclonal Rat Anti-Mouse IgD | Biolegend | 405723 | (dil 1/400) |
| Antibody                         | Monoclonal Rat Anti-Mouse IgD | Biolegend | 405710 | (dil 1/400) |
| Antibody                         | Monoclonal Rat Anti-Mouse IgG2b | Biolegend | 406708 | (dil 1/25) |
| Antibody                         | Monoclonal Rat Anti-Mouse IgM | Biolegend | 406512 | (dil 1/100) |
| Recombinant DNA reagent          | VDJ_H \text{NIMP23}^{\text{NIMP23}} \text{anti-MSP1}_{21} \text{variable region coding exon containing the Leader-V segment intron from gDNA of the NIMP23 hybridoma} | PMID: 7141700 | -- | Produced in Jean Langhorne’s lab |

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| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|------------------------------------|-------------|---------------------|-------------|------------------------|
| Recombinant DNA reagent            | C57Bl/6 IgH HEL variable region knock-in construct | PMID: 12668643 | – | Donated by Robert Brink of the Garvan Institute of Medical Research, New South Wales, Australia |
| Peptide, recombinant protein       | MSp121      | PMID: 11254580 | – | Produced in Jean Langhorne’s lab |
| Commercial assay or kit            | EZ-Link Suflo-NHS-LC-Biotinylation Kit | Thermo Scientific | 21435 | |
| Commercial assay or kit            | RiboPure RNA Purification Kit | Invitrogen | AM1924 | |
| Commercial assay or kit            | Qubit 1X dsDNA HS Assay Kit | Invitrogen | Q33231 | |
| Commercial assay or kit            | SMART-Seq v4 Ultra Low Input RNA Kit for Sequencing | Takara | 634889 | |
| Commercial assay or kit            | Ovation Ultralow Library System V2 | Nugen | 0344–32 | |
| Commercial assay or kit            | LIVE/DEAD Fixable Aqua Dead Cell Stain Kit | Invitrogen | L34957 | |
| Commercial assay or kit            | LIVE/DEAD Fixable Blue Dead Cell Stain Kit | Invitrogen | L23105 | |
| Chemical compound, drug            | Streptavidin-R-Phycoerythrin | Prozyme | PJRS25 | |
| Chemical compound, drug            | Streptavidin-Allophycocyanin | Prozyme | PJ27S | |
| Chemical compound, drug            | TiterMax Gold Adjuvant | Merck (formerly Sigma-Aldrich) | T2684-1ML | |
| Chemical compound, drug            | R848 (Resiquimod) | Invivogen | tlr1-r848 | |
| Chemical compound, drug            | TRI Reagent Solution | Invitrogen | AM9738 | |
| Software, algorithm                | cutadapt v1.9.1 | doi:10.14806/ej.17.1.200 | | |
| Software, algorithm                | RSEM v1.2.31 | doi:10.1186/1471-2105-12-323 | | |
| Software, algorithm                | STAR v2.5.1b | doi:10.1093/bioinformatics/bts635 | | |
| Software, algorithm                | DESeq2 | doi:10.1186/s13059-014-0550-8 | | |
| Software, algorithm                | R v3.4.0 | other | | |
| Software, algorithm                | Bioconductor v3.5 | other | | |
| Software, algorithm                | Broad’s Gene Set Enrichment Analysis (GSEA) | other | | |

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Continued

| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|--------------------|-------------|------------------------|
| Software, algorithm               | FlowJo version 9.6 or higher | Tree Star |             |                        |
| Software, algorithm               | Cytofkit | doi:10.1371/journal.pcbi.1005112.s009 |             |                        |
| Software, algorithm               | Prism v6 | GraphPad           |             |                        |

**Mice**

5-12 week-old female mice were used for experiments. C57BL/6J, C57BL/6.SJL-Ptprc<sup>a</sup> (CD45.1 congenic), Rag2<sup>-/-</sup>.C57BL/6.SJL-Ptprc<sup>c</sup> (CD45.1 congenic) and BALB/c mouse strains were bred in the specific pathogen-free facilities of the MRC National Institute for Medical Research and The Francis Crick Institute, and were housed conventionally with sterile bedding, food and irradiated water. Room temperature was 22°C with a 12 hr light/dark cycle; food and water were provided ad libitum. The study was carried out in accordance with the UK Animals (Scientific Procedures) Act 1986 (Home Office license 80/2538 and 70/8326), was approved by the MRC National Institute for Medical Research Ethical Committee and was approved by The Francis Crick Institute Ethical Committee.

To produce MSP1<sub>21</sub>-specific B cell knock-in mice capable of undergoing class switch recombination on the C57BL/6J genetic background, the VDJ<sub>H</sub>NIMP23 anti-MSP1<sub>21</sub> variable region coding exon containing the Leader-V segment intron from gDNA of the NIMP23 hybridoma (Boyle et al., 1982) was inserted by homologous recombination into the 5’ end of the endogenous IgH locus (Taki et al., 1993) (Figure 1—figure supplement 1A–C). The VDJ<sub>H</sub>NIMP23 anti-MSP1<sub>21</sub> variable region coding exon was inserted into a previously described IgH targeting construct, replacing the anti-HEL heavy chain variable region coding exon that was already in it (Phan et al., 2003) (Figure 1—figure supplement 1D). The final targeting construct included a loxP-flanked neomycin resistance cassette in reverse transcriptional orientation to the IgH locus, located immediately 5’ to the rearranged VDJ<sub>H</sub>NIMP23 variable region and its associated promoter (Figure 1—figure supplement 1D). Electroporation of C57BL/6N-derived PRX embryonic stem cells with the targeting construct and selection of homologous recombinant clones was performed using standard techniques by PolyGene AG (Switzerland). One targeted ES clone was used for production of chimeric mice using standard techniques at the Biological Research Facilities of the MRC National Institute for Medical Research, London, UK. Male chimeric mice were crossed to C57BL/6J females and progeny carrying the IgH<sup>NIMP23neo</sup> allele were crossed to PC3Cre mice (O’Gorman et al., 1997) to delete the neo<sup>r</sup> gene in the germline and generate mice carrying the IgH<sup>NIMP23</sup> allele (Figure 1—figure supplement 1D). The IgH<sup>NIMP23</sup>/+ strain was maintained by backcrossing for at least 10 generations to C57BL/6J mice.

**Mixed bone marrow chimeras**

Femurs and tibias were excised from female mice and cleaned of flesh using forceps and scalpel, and BM was obtained by flushing out with IMDM supplemented with 2 mM L-glutamine, 0.5 mM sodium pyruvate, 100 U penicillin, 100 mg streptomycin, 6 mM Hepes buffer, and 50 mM 2-ME (Gibco, Invitrogen), using a syringe with a needle. Thereafter, single BM cell suspensions were obtained by mashing through a 70 μm filter mesh, further sieved through 40 μm filter mesh and washed once. Live cells were resuspended in sodium chloride solution 0.9% (Sigma) at 4 × 10<sup>6</sup> cells/200 μl. Rag2<sup>-/-</sup>.C57BL/6.SJL-Ptprc<sup>a</sup> mice were sub-lethally irradiated (5Gy) using a [137Cs] source at 4 × 10<sup>6</sup> cells/200 μl. Rag2<sup>-/-</sup>.C57BL/6.SJL-Ptprc<sup>a</sup> mice were sub-lethally irradiated (5Gy) using a [137Cs] source and reconstituted less than 24 hr after irradiation by i.v. injections of a 10% IgH<sup>NIMP23</sup>/:90% C57BL/6.SJL-Ptprc<sup>a</sup> combination of donor BM cells. Recipient mice were maintained on acidified drinking water and analyzed for reconstitution after 6–8 weeks.

**Plasmodium chabaudi infection**

*Plasmodium chabaudi chabaudi* AS was transmitted by Anopheles stephensi mosquitoes, strain SD500, as described elsewhere (Spence et al., 2012). Briefly, C57BL/6J mice were injected i.p. with 10<sup>5</sup> P. chabaudi-infected red blood cells and used to feed mosquitoes two weeks after the injection.
Two weeks after mosquito feeding/infection, each experimental mouse was exposed to 20 infected mosquitoes for 30 min. Blood parasitemia in infected experimental mice was routinely monitored by thin blood smears.

**Immunizations**

Mice were immunized i.p. with a combination of 100 µg of MSP1$_{21}$ (Quin and Langhorne, 2001) and 50 µl of Titermax Gold emulsion (Sigma), or a combination of 50 µg of MSP1$_{21}$ and 50 µg of R848 (Invivogen).

**Flow cytometry and cell sorting**

Spleens, lymph nodes and bone marrows were dissected and single cell suspensions were obtained by mashing the organs through a 70 µm filter mesh in HBSS, 6 mM Hepes buffer (Gibco, Invitrogen). After removal of red blood cells from spleens and bone marrows by treatment with lysing buffer (Sigma), the remaining cells were resuspended in complete Iscove’s Modified Dulbecco’s Medium (IMDM supplemented with 10% FBS Serum Gold (PAA Laboratories, GE Healthcare), 2 mM L-glutamine, 0.5 mM sodium pyruvate, 100U penicillin, 6 mM Hepes buffer, and 50 mM 2-ME (all from Gibco, Invitrogen)) and viable cells were counted using trypan blue (Sigma) exclusion and a hemocytometer. Cells were then resuspended in PBS and incubated with APC- or PE-labelled MSP1$_{21}$ fluorescent probes and/or different combinations of fluochrome-conjugated antibodies (key resources table), and either acquired after two washes with PBS, or fixed with 2% paraformaldehyde and stored in staining buffer at 4°C until acquisition.

The APC and PE MSP1$_{21}$ fluorescent probes were produced as previously described (Krishnamurty et al., 2016; Taylor et al., 2012). Briefly, purified, modified MSP1$_{21}$ (Quin and Langhorne, 2001) was biotinylated using an EZ-link Sulfo-NHS-LC- Biotinylation kit (Thermo Fisher Scientific) using a 1:1 ratio of biotin to protein, and loaded onto Streptavidin-APC conjugated or Phycolink Streptavidin-R-PE conjugated (ProZyme) in a 6:1 ratio of MSP1$_{21}$:Streptavidin-fluorochrome.

Cell sorting was performed on a MoFlo XDP (Beckman Coulter) or a BD FACSAria Fusion (BD Biosciences) and the target cell populations were directly dispensed into TRIreagent (Ambion) and stored at -80°C until RNA isolation. Purity checks were routinely performed for all assays by sorting aliquots of cells into PBS containing 2% FCS and reacquiring them on the cell sorter.

Dead cells were routinely excluded from the analysis by staining with LIVE/DEAD Fixable Aqua or Blue stain (Invitrogen). Singlets were selected based on FSC-A vs FSC-H and further based on SSC-A vs SSC-H. ‘Fluorescence minus one’ (FMO) controls were routinely used to verify correct compensation and to set the thresholds for positive/negative events. Analysis was performed with FlowJo software version 9.6 or higher (Tree Star).

PhenoGraph and t-distributed stochastic neighbor embedding (t-SNE) were combined to analyze multiparameter flow cytometry data using the Cytofkit package (Chen et al., 2016). t-SNE renders high-dimensional single-cell data based on similarities into only two dimensions, and thus helps visualize multiparameter data (van der Maaten, 2008). PhenoGraph (Levine et al., 2015) allows partitioning of high-dimensional single-cell data into phenotypically coherent subpopulations (i.e. clusters). The relatedness of the cell clusters identified by PhenoGraph was inferred using Isomap (Cytofkit package), in which related clusters/subsets can be visualised close to each other.

**RNA isolation, sequencing and data analysis**

Total RNA from 1–5 × 10$^4$ cells sorted into TRIreagent (Ambion) was isolated using the Ribopure kit (Ambion). Concentration of purified RNA was determined by Qubit fluorometric quantitation using the HS assay kit (ThermoFisher Scientific), and the quality analyzed with a 2100 Bioanalyzer (Agilent). Samples with a RIN score above 8.50 were used for the next steps. cDNA was generated from total RNA with the SMART-Seq v4 Ultra Low Input RNA Kit (Takara Bio USA). Next-generation sequencing libraries were produced with the Ovation Ultralow System V2 (Nugen), and run as PE100 on a HiSeq 4000 sequencer (Illumina). GEO accession: GSE115155.

For bioinformatics analysis, paired-end sequence reads were adapter and quality trimmed using cutadapt v1.9.1 (Martin, 2011) with the following non-default settings: ‘-a AGATCGGAAGAGC -A AGATCGGAAGAGC –minimum-length 30 -q 20,20’. Gene-level abundance estimates were generated from the trimmed reads using RSEM v1.2.31 (Li and Dewey, 2011) running STAR v2.5.1b
(Dobin et al., 2013) with default settings, aligned against the Mus musculus Ensembl release 89 transcriptome (mm10). All further analysis was conducted using the DESeq2 (Love et al., 2014) package from Bioconductor v3.5 run in R v3.4.0. The expected counts were imported and rounded to integers to generate a counts matrix. Differential expression between phenotype groups was assessed using the DESeq function with default settings. In the case of comparisons of different MSP121-specific B cell subsets obtained from the same experimental mouse, an additional mouse factor was added to the design formula to accommodate the paired nature of the data. Significance was thresholded using an FDR ≤ 0.01. PCA analysis was conducted using DESeq’s plot PCA function with the regularized log (rlog) transformed count data. Heat maps were generated using the regularized log (rlog) transformed count data, scaled per gene using a z-score. Mouse homologues to genes previously associated with human AMB were selected (Supplementary file 1), and those showing significant differential expression on MSP121-specific AMB were used to produce separate heat maps split by functional annotation (Figure 3). The GSEA pre-ranked function from the Broad’s Gene Set Enrichment Analysis (GSEA) (Subramanian et al., 2005) suite was used to assess significant enrichment of MSigDB’s C2 Reactome gene sets associated with differential expression between cell types. The function was run using a list of genes ranked for differential expression using DESeq2’s Wald test statistic with default settings except for: collapse dataset to gene = false enrichment statistic = classic

Statistical analysis
Statistical analysis was performed using Mann Whitney U test, Kruskal-Wallis test followed by Dunn’s multiple comparisons test, or Two-Way ANOVA followed by Dunnett’s multiple comparisons test on Prism software version 6 (GraphPad). p<0.05 was accepted as a statistically significant difference.

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Animal experimentation: The study was carried out in accordance with the UK Animals (Scientific Procedures) Act 1986, Home Office license 80/2538 and 70/8326. Was approved by the MRC National Institute for Medical Research Ethical Committee, The Francis Crick Institute Ethical Committee, and further approved by the Home Office of the UK upon granting of the HO license.

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Additional files
Supplementary files
• Supplementary file 1. Mouse homologues to human genes previously described to be either up (↑) or down (↓) regulated in human atypical memory B cells (AMB).
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• Supplementary file 2. List of top 50 Reactome gene sets yielding the highest normalized enrichment score (NES) by GSEA.
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Data availability
RNAseq transcriptome data have been deposited in GEO under accession code GSE115155.

The following dataset was generated:

| Author(s)                           | Year | Dataset title                                                                 | Dataset URL                                                                 | Database and Identifier                  |
|-------------------------------------|------|-------------------------------------------------------------------------------|----------------------------------------------------------------------------|------------------------------------------|
| Pérez-Mazliah D, Gardner PJ,       | 2018 | Plasmodium-specific atypical memory B cells are short-lived activated B cells | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE115155               | NCBI Gene Expression Omnibus, GSE115155  |
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