Identification of MicroRNAs Regulating the Developmental Pathways of Bone Marrow Derived Mast Cells

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Abstract

Background: MicroRNAs (miRNAs) play important roles in leukocyte differentiation, although those utilised for specific programs and key functions remain incompletely characterised. As a global approach to gain insights into the potential regulatory role of miRNA in mast cell differentiation we characterised expression in BM cultures from the initiation of differentiation. In cultures enriched in differentiating mast cells we characterised miRNA expression and identified miRNA targeting the miRNA of putative factors involved in differentiation pathways and cellular identity. Detailed pathway analysis identified a unique miRNA network that is intimately linked to the mast cell differentiation program.

Methodology/Principal Findings: We identified 86 unique miRNAs with expression patterns that were up- or down-regulated at 5-fold or more during bone marrow derived mast cells (BMMMC) development. By employing TargetScan and MeSH databases, we identified 524 transcripts involved in 30 canonical pathways as potentially regulated by these specific 86 miRNAs. Furthermore, by applying miRanda and IPA analyses, we predict that 7 specific miRNAs of this group are directly associated with the expression of c-Kit and FcεRIα and likewise, that 18 miRNAs promote expression of Mitf, GATA1 and c/EBPα, three core transcription factors that direct mast cell differentiation. Furthermore, we have identified 11 miRNAs that may regulate the expression of STAT5, -5a/b, GATA2 and GATA3 during differentiation, along with 13 miRNAs that target transcripts encoding Ndst2, mMCP4 and mMCP6 and thus may regulate biosynthesis of mast cell secretory mediators.

Conclusions/Significance: This investigation characterises changes in miRNA expression in whole BM cultures during the differentiation of mast cells and predicts functional links between miRNAs and their target miRNAs for the regulation of development. This information provides an important resource for further investigations of the contributions of miRNAs to mast cell differentiation and function.

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Introduction

Mast cells (MCs) are vital innate immune cells that are well-known for their roles in the pathogenesis of immunoglobulin E (IgE)-mediated disorders [1], while aberrant generation of MCs can lead to disorders such as urticaria pigmentosa, systemic mastocytosis and mast cell activation syndrome [2–4]. There is also increasing recognition of the importance of MCs to both innate and adaptive host defence against bacterial and parasitic infection [5,6]. Thus, gaining a better understanding of pathways that control MC differentiation will provide important insights into the developmental programs of MCs in both health and in disease states.

MCs are derived from pluripotent progenitors that originate in bone marrow and that mature in most tissues but are particularly enriched in the skin, tongue and in the mucosal layer of lung and gastrointestinal tracts, under the control of specific growth and differentiation factors [7]. Expansion of MCs predominantly and synergistically relies on stem cell factor (SCF) and interleukin (IL)-3 [8]; both human and murine MCs express receptors for these two cytokines [9,10]. Interestingly, IL-3 alone cannot generate MCs, but this cytokine greatly enhances the numbers of differentiated mast cells in tissue culture and in vivo [10,11]. By contrast, mice with SCF receptor (c-Kit) mutations [Kitmsh/msh] and Kitw-sh/w-sh mice) have drastically reduced c-Kit/SCF signaling and demonstrate a profound reduction in MCs [12,13]. Likewise, clinical
evidence suggests that an activating mutation in the c-Kit gene is strongly associated with systemic mastocytosis [14]. Although SCF and IL-3 drive the development of MCs, the transcription factors and pathways that specifically control differentiation have not been fully characterised. Currently, microphthalmia-associated transcription factor (Mitf) is considered as a central mediator of MC differentiation; Mitf is highly expressed in both progenitor and fully differentiated MCs, while notably absent in basophils [15]. Furthermore, accumulating evidence suggests that the interplay between four co-re-regulatory
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Methods

Animals
Specific pathogen free BALB/c mice (6-8 weeks) were obtained from the animal services unit of the University of Newcastle. All experiments were performed with approval from the Animal Ethics Committee of the University of Newcastle (Permit Number: A-2010-136). All surgery was performed under sodium pentobarbital anesthesia, and all efforts were made to minimize suffering.

Culture and identification of bone marrow derived mast cells
Bone marrow derived mast cells (BMMCs) were differentiated and examined as previously described, with slight modifications [23]. Briefly, mouse femurs were flushed with 5 ml ice cold HBSS through a 70 μm cell strainer. After lysis of red blood cells and washing with PBS, BMMCs were cultured for 6 weeks in RPMI 1640 medium (Invitrogen) supplemented with 10% FCS, 4 mM glutamine, 100 U/ml penicillin, 100 μg/ml streptomycin, 25 mM hydroxyethyl piperazineethanesulfonic acid, 1 mM sodium pyruvate, and 50 μM 2-mercaptoethanol, 15 ng/ml SCF (Peprotech) and 30 ng/ml IL-3 (Peprotech) at a starting concentration of 1×10^6 cells/ml and cultured at 37°C in a humidified atmosphere of 5% CO2 and 95% air. Culture medium was replaced twice per week for 6 weeks and the cell concentration was maintained at 1×10^6 cells/ml. 2.5×10^7 cultured cells were harvested every week for miRNA microarray and qPCR. The purity of BMMCs was determined by using toluidine blue staining of cytospins (1×10^6 BMMCs per slide) and flow cytometry (See Method S1).

MicroRNA microarray
Total RNA was extracted from bone marrow cells and BMMC cultures from week 2 to week 6 using TRizol reagents (Life Technologies), and miRNA microarray was performed as previously reported [24]. Briefly, the Agilent Agilent spike-In control was added to 100 ng RNA, which was dephosphorylated by incubating the samples at 37°C for 30 minutes following by ligation of Cy3 using the Complete Labelling and Hybridisation c-Kit (Agilent). Following ligation and drying, the Cy3-labelled RNA samples were hybridized for 20 h at 55°C to Agilent 8x15K mouse microRNA array slides (AMADID 21828), which included 627 mouse miRNA and 39 mouse viral miRNA from the Sanger database 12.0. After washing with Agilent gene expression wash buffers, the hybridized microarrays were scanned on a High Resolution C scanner (Agilent). Data were extracted from scanned microarrays using Feature Extraction software (version 10.7.3.1). The fluorescence index of each miRNA at different time-points was further normalized to that of the respective miRNAs in control group (isolated bone marrow cells). The normalized microarray data were managed and analyzed by GeneSpring (Agilent). MicroArray data have been deposited into ArrayExpress (http://www.ebi.ac.uk/arrayexpress/). The accession number is E-MTAB-2443.

Quantification of microRNA and Messenger RNA
Quantitative polymerase chain reactions (qPCR) were performed as previously described (also See Method S1) [24]. Primer sequences are shown in Table S1.

MicroRNA target analysis
For prediction of target genes of differentially expressed miRNAs, we first used TargetScan 6.1 (http://www.targetscan.org/) to identify potential targets. MeSH database (http://www.ncbi.nlm.nih.gov/mesh/meshhome.html) was then employed to identify the molecules relevant to MC biology by exact syntax matching. MiRanda (http://www.microrna.org/) was also used to refine the predicted targets. Ingenuity Pathways Analysis (Ingenuity Systems, Redwood City, CA) software was employed so as to define canonical signaling pathways containing the miRNA-associated MC-relevant molecules, and to establish the connection between miRNAs and their respective predicted targets.

Statistical analysis
An initial one-way ANOVA was followed by appropriate comparisons to test for differences between means of groups. Values are reported as the means ± SEM for each experimental
group. The number of samples at each time-point ranged from 4 to 6. Differences in means were considered significant if \( p < 0.05 \).

**Results**

**Generation of BMMC**

Isolated bone marrow cells from BALB/c mice were cultured for 6 weeks in the presence of SCF and IL-3. From week 2 to week 6, the percentage of BMMC (c-Kit^+ FcεRI^+) in the culture was determined first by flow cytometry. The morphological features, the percentages and numbers of BMMCs were also determined by toluidine blue staining of cytospin preparations. The percentage of c-Kit^+ FcεRI^+ cells increased from 4.2\% to 92.7\% in BMMC culture at week 6 (Fig. 1A). We observed a parallel increase in toluidine-blue positive cells (5.34 ± 1.9\% to 99.0 ± 0.03\%; 2.70 ± 0.21 × 10^5 to 36.5 ± 3.2 × 10^5 cells/mL; Figs. 1B and C, respectively.)

**Distinct miRNA expression profile during BMMC development**

Given the broad involvement of miRNAs in regulating post-transcriptional gene expression, we proceeded to determine the miRNA profile of BMMC cultures from week 2 to week 6. Total RNA was isolated from homogenised cells and screened as...

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**Figure 3. Potential molecules and top canonical pathways that were predicted and targeted by the miRNAs identified as differentially regulated during BMMC differentiation.** (A) Target prediction by TargetScan database (http://www.targetscan.org/) was established on sequence data complementarity to target 3’ UTR sites. Target molecules, associated with MC biology, were identified by exact syntax matching in the MeSH database (http://www.nlm.nih.gov/MeSH/MeSHhome.html). (B) Top 30 canonical pathways that include selected 524 molecules as identified by IPA. The significance of association between selected genes and canonical pathway was evaluated by a right-tailed Fisher’s exact test (grey bars, upper y-axis). Ratios referring to the proportion of selected genes from a pathway related to the total number of molecules that make up that particular pathway were also displayed (line graph, bottom y-axis). doi:10.1371/journal.pone.0098139.g003
described in Methods. We identified eighty-six (86) miRNAs characterised with 5-fold differential expression during BMMC development; among these, we document 45 miRNAs with increased expression and 41 with decreased expression, respectively (Fig. 2 A).

Detailed information describing these miRNAs is included in Table S2 in File S1. We validated the differential expression of six (6) of the 86 miRNAs with qPCR (Fig. 2 B). Among the miRNAs selected for validation, miR-223 has a central role in myeloid cell development [25]. The other miRNAs have been linked to cell development, inflammation and disease [26–30]. The expression profiles defined for these six miRNAs by qPCR were consistent with the expression patterns documented on the microarray.

Analysis of BMMC related molecules and canonical pathways potentially regulated by the miRNAs

In an effort to elucidate targets and pathways regulated by the 86 differentially-expressed miRNAs, we proceeded to identify...
putative mRNA targets using TargetScan (version 6.1). TargetScan is a search engine that predicts targets for miRNA-mRNA interactions in eukaryotes; the algorithms search for 8- and 7-mer sites located in the 3′-UTR regions of mRNAs which share sequence homology with query miRNA sequences [31]; predictions are then ranked by their probability of targeting the mRNA transcript [32]. We used a 95% percentile cutoff to predict the potential targets. With this method, we identified 5924 mRNA transcripts that could be targeted by the 86 miRNAs (Fig. 3 A). The target mRNAs were further refined by searching the MeSH database, in an effort to link gene expression data with known MC-related pathways, diseases and phenotypes. There were 1418 mRNAs identified by MeSH database as containing the exact terms “mast cell, SCF, IL-3 and tryptase” with a link to at least one PubMed-affiliated reference. We then plotted differences in distribution of mRNAs predicted by TargetScan and MeSH databases, and found that, of the original 5924 mRNA transcripts, we could focus on 524 that were targeted by both search methods (Table S3 in File S1).

These 524 MC mRNAs were then organized and classified with the assistance of Ingenuity Pathways Analysis (IPA); the top 30 canonical pathways were listed (Fig. 3 B; Table S4 in File S1). Two-hundred and three (203) of the aforementioned molecules (38.7% of the 524 MC-associated transcripts) were included within these top 30 canonical pathways. Although many of the pathways are commonly involved in cell death and survival or proinflammatory activity, these pathways also include those that contribute to mast cell differentiation and function. For example, Phosphatase and Tensin Homolog (PTEN) regulates the proliferation MC; PTEN deficiency results in a mastocytosis-like proliferative disease [33]. Likewise, activation of Glucocorticoid Receptor signaling suppresses both proliferation and degranulation of MCs [34], and PI3/AKT signaling is critical for the maturation and pro-inflammatory factor production by MC [35]. Pathways such as Growth Hormone, IGF-1, Apoptosis and Role of JAK2 in Hormone-Like Cytokine may regulate MC survival and death [8]. FLT3 Signaling in Hematopoietic Progenitor Cells, Erythropoietin, IL-6 and ErbB signaling may play central roles in differentiation of haematopoietic progenitor cells to MC specific lineage [36]. Well-known inflammatory pathways such as Acute Phase Response and NF-kB likewise regulate the production of pro-inflammatory factors in MCs, and NF-kB can also modulate MC proliferation [37,38]. TREM1 may play a central role in MC responses to bacterial invasion [37]. Interestingly, activation of MCs has long been recognized as an important component of the pathogenesis of rheumatoid arthritis [39]; the two signaling pathways, Role of Osteoblast/Osteoclast and Chondrocytes in Rheumatoid Arthritis and Role of Macrophages/Fibroblast and Endothelial Cell in Rheumatoid Arthritis, may feature components of this role. Taken together, these data suggest important roles for miRNAs in the regulation of trafficking, biological function and cell death/cell survival of BMMCs.

Multiple miRNAs are linked to the expression of c-Kit and FceRI but not IL-3Rα.

IL-3Rα, c-Kit and FceRI are critical signature receptors of MCs; we examined the expression of these molecules in BMMC cultures from week 2 to week 6 by qPCR (Fig. 4 A). Relative expression of transcripts encoding IL-3Rα underwent a small but statistically significant increase between week 2 and week 5. By contrast, expression levels of c-Kit and FceRIα increased markedly (greater than 40-fold over baseline by week 6). We used IPA and the miRanda database in order to determine whether any of the 86 differentially expressed miRNAs (Fig. 2 A) might have binding sequences that could target the 3′-UTRs of any of these specific transcripts. Interestingly, none of the aforementioned 86 miRNAs has any clear potential to regulate IL-3Rα. However, the c-Kit transcript is linked to 6 specific miRNAs, of which 3 (miR -218, -505 and -542-3p) undergo an increase and 3 (miR -130a, -223 and -421) a decrease (Fig. 4 B and C) during BMMC differentiation in culture. The transcript encoding FcεRIα may be targeted by miR-363. The 3′-UTR binding sites of these miRNAs are included in Table S5 in File S2.

Critical MC transcriptional regulators targeted by distinct sets of miRNAs.

The fate of BMMCs is critically determined by a coordinated action by transcription factors Mitf, GATA1, c/EBPα and PU.1 [17,18]. We evaluated the differential expression of these factors by qPCR and correlated these patterns to those of the 86 differentially expressed miRNAs (Fig. 2 A). As previously reported [15,16], expression levels of Mitf and GATA1 were significantly elevated during differentiation of BMMCs while that of c/EBPα decreased dramatically (Fig. 5 A). PU.1 expression did not change, although it was highly expressed overall in BMMC culture (data not shown). Using IPA and miRanda, we identified three sets of miRNAs that have to potential to regulate the expression of these four MC transcriptional regulators (Fig. 5 B). We correlated the expression levels of specific miRNAs to Mitf (miR -181d, -207, -298, -342-5p, -710 and -1896), GATA1 (let-7e, miR -196b, 139-5p, -200b and -290-5p) and c/EBPα (miR -129-3p, -218, -330, -363, -672, 1894-3p and 671-5p), but interestingly, found no miRNAs with the capacity to target PU.1 (Fig. 5 C). The 3′-UTR binding sites of miRNAs that target all three of these miRNAs are shown in Table S6 in File S2. These findings likewise suggest that miRNAs have the potential for extensive and critical involvement in the regulation of MC differentiation.
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A. 

|   | Stat3  |   | Stat5a |   | Stat5b |
|---|--------|---|--------|---|--------|
| BM w2 w4 w6 | Fold changes | BM w2 w4 w6 | Fold changes | BM w2 w4 w6 | Fold changes |

B. 

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miR-874 → miR-23b
miR-330 → Stat3 → miR-125b-5p → Stat5b → miR-342-3p
miR-34a → miR-132
miR-135a → miR-290-5p → Gata3 → Gata2 → miR-200b → miR-363
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C. 

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miR-330 → 200
miR-874 → 125b-5p
miR-125b-5p → 342-3p
miR-34a → 132
miR-135a → 290-5p
miR-290-5p → 200b
miR-200b → 363
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myeloid progenitors [1,16–18]. We first determined the expression levels of STAT3, STAT5a, STAT5b, GATA2 and GATA3 during BMMC culture (Fig. 6A). All of these factors responded to culture conditions with increased expression from week 2 to week 6. Interestingly, the miRNAs that targeted these mRNA transcripts (Fig. 6B), further refined by IPA and miRanda analysis, mostly overlapped with those identified in Fig. 5B and Fig. 6B. The interactions between the 3′-UTRs of these five mRNAs and their associated miRNAs are presented in Table S7 in File S2. For STAT3, the two of the three associated miRNAs (miR −330 and −874) responded with increased expression and one (miR−125b-3p) decreased; for STAT5b, two miRNAs (miR−125b-5p and −342-3p) responded with decreased expression and one (miR−23b) increased (Fig. 6C). Decreased expression of miR−342-3p was also associated with increased levels of STAT5a, and we found that one factor, miRNA-132, may concurrently target STAT5b, GATA2 and GATA3. Notably, there is considerable overlap between the miRNAs that regulate the three core transcription factors (Fig. 5), and the aforementioned STATs and GATA2/GATA3 regulators (Fig. 6). The distinct role of these miRNAs in fine-tuning the expression of these molecules in MC differentiation remains to be explored.

Association between miRNAs expression and major functional transcripts of BMMC

MC granules contain substantial amounts of heparin, histamine, MC-specific proteases (tryptase and chymase) and other unique biological factors [7]. N-deacetylase/N-sulfotransferase (Ndst) 2 is a crucial enzyme in the biosynthetic pathway of heparin and its expression is strongly linked to the process of MC maturation [40]. Likewise, histamine is synthesized by an enzymatic reaction catalyzed by L-histidine decarboxylase (HDC), while MC proteases are each encoded by the unique genes, mast cell protease (mMCP) −2, −4, −5, −6 and −7 [7]. We determined that transcripts encoding Ndst2, mMCp6 and mMCp7 were significantly up-regulated from week 2 to week 6, whereas the expression of myeloperoxidase (MPO, predominantly expressed in neutrophils) decreased to nearly undetectable levels at wk 6 (Fig. 7A). HDC was constantly expressed at high levels over the same interval (unpublished observation). Mouse MCP4 increased marginally at week 2 but was down-regulated to 21% of initial levels by week 6 (Fig. 7A). Ndst2 expression was linked to that of three miRNAs (let-7e, miR −211 and −207) that showed increased expression and three miRNAs (miR −153a, −671-5p and −1894-3p) with decreased expression (Fig. 7B and C). While mMCp4 was associated with the increased expression of 5 miRNAs (miR −132, −330, 501-3p, 542-3p and −672) and the decreased expression of miR−1894-3p (Fig. 7B and C), MPO and mMCp6, were targeted by miR−207 and miR−705 which had increased and decreased expression, respectively. The interaction between 3′-UTR of the four mRNAs and their associated miRNAs are presented in Table S8 in File S2. Interestingly, the sequence of miR−1893-3p suggests that it is likely to target both Ndst2 and mMCp4.

Discussion

Previous studies have identified several miRNAs, including miR−132, miR−146a and miR−221, that play roles in regulating MC apoptosis, cell cycle regulation, growth and degranulation [41–43]. Monticelli and colleagues have used 181-miRNAs-Array to profile matured BMMC and identified 15 of them with detectable changes. However, their method can not efficiently discriminate between close miRNA paralogs [44]. As a more recent example, Lee and colleagues have shown that decreased expression of both miR−539 and miR−381 in association with c-Kit signaling promotes expression of the transcription factor, Mitf, which leads to MC proliferation [45]. Our present study was undertaken in order to provide a comprehensive examination of mRNA temporal expression during differentiation of BMMC in culture starting within the initiation of differentiation in cellular milieu of the BM, with more extensive and gene-specific Agilent Array containing 627 mouse miRNA and 29 mouse viral miRNA. Then by using detailed pathway analysis programs we have linked miRNAs expression in cultures enriched in differentiating mast cells with the regulation of transcripts that play critical roles in mast cell development. Toward this end, we have established a BMMC ex vivo culture system and we have determined the expression of characteristic receptors, functional factors and critical transcription regulators during the development. The profile of expressed miRNAs was determined over time and correlated to expression of molecules known to be crucial to MC development by stringent statistical and validated pathways analyses. Via these methods, we documented an original MC expression signature that included 86 independent miRNAs that were differentially regulated five-fold or more over baseline levels during the development of BMMC between week 2 and week 6.

MiRNAs play critical roles in maintaining gene expression at appropriate levels and in fine-tuning a wide range of biological and disease processes [20–22]. However, it remains difficult to associate miRNAs with their targets either directly or experimentally and likewise to predict specific biological functions. For example, Siemens and colleagues have determined that c-Kit is a direct target of miR−34a by using several human colorectal cancer cell lines [46]. Although miR−34a has two continuous 5-mer binding sites at the 3′-UTR of human c-Kit, it may bind weakly to the respective region of mouse c-Kit as there are only two discontinuous 5-mer seed sequences. This suggests that miR−34a is unlikely to directly regulate the expression of c-Kit in mouse. Interestingly, Shin and colleagues have identified that miR−34a contributes to BMMC apoptosis by targeting Bcl-2 [47]. These observations may show that the potential targets of miRNAs can be rather distinctive and these small regulatory molecules play fairly distant regulatory roles in biological development and the progression of diseases in different species.

Here, we have employed a combinatorial approach, and we searched for targets for the 86 differentially-regulated miRNAs initially based on algorithms from TargetScan and further filtered the potential targets by searching MeSH database for published links to MC biology. From there, our findings were refined and categorized into networks that associated miRNAs with critical MC factors using miRanda and IPA. With this approach, we have
Figure 7. Expression of Ndst2, MPO, mMCP4 and mMCP6 correlated to the expression of miRNAs that potentially target these transcripts. (A) Expression levels of Ndst2, MPO, mMCP4 and mMCP6 were determined by qPCR. (B) Potential miRNAs targeting the 3′-UTR of Ndst2, MPO, mMCP4 and mMCP6 were identified by TargetScan and Miranda database and IPA ingenuity system. Blue represents decreased expression of miRNAs, whereas yellow is for increased expressed miRNAs. (C) The fold changes of potential regulating miRNAs were calculated based on the fluorescence index of each miRNA at different time-points, after normalization to that of the respective miRNAs in the control group (isolated bone marrow cells). Values are presented as mean ± SEM (n = 4–6), **P < 0.001 (vs. BM), *P < 0.01 (vs. BM). doi:10.1371/journal.pone.0098139.g007
identified 32 unique miRNAs that are most likely to be involved in regulating MC differentiation and function via targeting critical MC factors.

By cross-comparison with TargetScan and MeSH database, we generated a list of 524 transcripts that have the highest-likelihood of serving as targets for the 96 differentially-expressed miRNAs and linked to MC biology. By IPA, we showed that 39% of these molecules were included within the top 30 signal transduction pathways, among which were molecules controlling cell cycle, growth and death, and cell activation. For example, PTEN and PI3K/AKT signaling are closely paired and serve to balance one another in order to regulate cell proliferation. Impairment of PTEN signaling may lead to PI3K/AKT hyperreactivity, resulting reduced apoptosis and unchecked cell proliferation [33]. Furthermore, miRNA may also target other pathways such as growth hormone, IGF-1 and FLT3 signaling to control MC differentiation. JAK/STAT and NF-kB signal transduction pathways that are responsible for cell differentiation and are pro-inflammatory are also potentially targeted by the identified miRNAs.

By examining connectivity between predicted targets, we distinguished between closely-linked molecules that may operate as network hubs, with highly associated factors functioning as similar units. Among the 524 potential miRNA targets, c-Kit and FcεRIα are the well-known signature receptors of MC [1,9,10]. The expression of c-Kit and FcεRIα were increased almost 400-fold from day 0 to 6 weeks of culture [5, A]. These results are consistent with the greatly expanded c-Kit and FcεRIα+ BM population. Although SCF and IL-3 are used for 4 to 8 weeks in most ex vivo BMNC cultures [8–13,23], we have observed that none of 86 miRNAs target the IL-3Rα chain transcripts. However, transcripts for this receptor were only slightly increased during culture and levels were high prior to induction of differentiation indicating that this system was already functional (data not shown). Taken together, these results suggest that the IL-3 signaling pathway may be involved more broadly in myeloid differentiation and its contributions are not necessarily limited to MC maturation. By contrast, the interactions between SCF and c-Kit are essential for MC development; deleting c-Kit in a mouse model results in deficiency of mast cells and elimination of IgE-mediated responses [12,13]. Six miRNAs were closely associated to c-Kit expression during MC differentiation; miR−218, −421 and −542-3p recognize multiple sites in the 3′-UTR (Table S4 in File S1), suggesting that these miRNAs are critical for MC function. Furthermore, it appears that the expression of FcεRIα is only regulated by miR-363 from this dataset.

As per current models, cooperation between several transcriptional regulators determines the differentiation and function of mast cells [1,16–18]. Our results suggest that three core transcriptional factors, Mitf, GATA1 and c/EBPα are differentially regulated by distinct groups of miRNAs (Fig. 6). The level of PU.1, which is another core regulator, did not change during BMNC culture, but levels of transcript remained high throughout (data not shown). Of note, there are other transcriptional factors involved in MC differentiation, such as STAT3, STAT3α/b and GATA2/3 [1,16–18]. Indeed, appropriate activation of these transcriptional pathways in cell growth, differentiation, survival, chemotaxis and cytokine production is essential for MC growth and differentiation [1,16–18]. Unlike the discrete connection between the distinct groups of miRNAs and Mitf, GATA1 and c/EBPα, the relationship between the STAT and GATA transcripts and miRNAs is more difficult to define (Fig. 7). This is partly because these proteins are directly activated by SCF and IL-3, and partly because that the activation of these transcription factors is largely regulated by their phosphorylation, rather than directly by increased protein expression. Nevertheless, it is noteworthy that miR-342-3p -with almost 1000-fold decreased expression- still targets STAT5α, STAT3β and Mitf. Furthermore, miR-125b-5p, that was decreased approximately 200 fold, is associated with the increased expression of STAT3 and STAT5β transcripts. This suggests that the down-regulation of the two aforementioned miRNAs potentially play critical roles in MC differentiation, particularly in the early phases of lineage commitment. On the contrary, miR-132 was significantly increased, but its potential targets (e.g. STAT5α and GATA2/3) were also increased. These findings suggest that miR-132 acts as an important checkpoint to maintain the expression of these transcriptional regulators at appropriate levels. Interestingly, miR-132 may also control the expression of the mast cell protease, mMCP-4, suggesting the broader roles for this miRNA in the MC biology.

Finally, to explore the connections between miRNAs and MC secretory function, we examined associations between selected miRNAs and the 3′UTRs of major MC secretory mediators. Among the thirteen miRNAs identified, interestingly, three miRNAs (miR−211, −501-3p and −705) were only linked to Ndo1, MCP4 and MCP6, respectively (Fig. 7). Not only with selected MC secretory mediators, seven of aforementioned 13 miRNAs (let-7e, miR−207, −298, −330, −671-3p, −672 and −1894-3p) were also associated with three core MC transcriptional regulators (Mitf, GATA1 and c/EBPα) (Figs. 6 and 8). Collectively, our data suggest that the molecular mechanisms controlling MC differentiation and function are closely interwoven where sets of miRNA play central roles.

Taken together, we present a model of BMNC differentiation via which we can examine expression patterns of miRNAs. We have analysed these patterns and identified miRNAs that may play pivotal roles in driving MC maturation and regulating MC function. However, to understand how these selected miRNAs interact with their targets, functional studies with miRNA targeting, knock-in and knock-down models are required. Nevertheless, we are able to demonstrate that these selected miRNAs can be grouped as networks in order to guide bone marrow cells toward MC development in cooperation with cytokine signaling. Furthermore, there are also a number of intracellular signaling pathways that may be regulated by these miRNAs. Manipulating the expression of these miRNAs may promote further understanding of the underlying mechanisms that are critical for MC differentiation and function, and ultimately provide therapeutic targets for treating MC-associated diseases.

Supporting Information

Method S1 Toluidine blue staining, flow cytometry and miRNA/mRNA qPCR were described.

(DOCX)

File S1 This file includes: Table S2. Annotation of miRNAs with greater than 5 fold changes, as shown in Figure 2; Table S3. BMNC related molecules potentially targeted by the miRNAs; and Table S4. BMNC related canonical pathways potentially regulated by the miRNAs.

(XLS)

File S2 This file includes: Table S5: Potential binding sites between c-Kit and FcεRIα and their respective miRNAs; Table S6: Potential binding sites between Mitf, c/EBPα and Gata1, and their respective miRNAs; Table S7: Potential binding sites between Stat3, Stat5α, Stat5β, Gata3 and Gata2, and their respective miRNAs; and
Table S8: Potential binding sites between Ndst2, mMCP4, mMCP6 and MPO, and their respective miRNAs.

Table S1 Primer sequence for determining mRNA levels by quantitative PCR.

Author Contributions
Conceived and designed the experiments: PSF MY. Performed the experiments: YX FE. Analyzed the data: YX FE IGY HFR PSF MY. Wrote the paper: HFR PSF MY.