Compartmentalization of innate immune responses in the central nervous system during cryptococcal meningitis/HIV coinfection


Published in:
AIDS

DOI:
10.1097/QAD.0000000000000200

Document Version
Peer reviewed version

Link to publication in the UWA Research Repository

General rights
Copyright owners retain the copyright for their material stored in the UWA Research Repository. The University grants no end-user rights beyond those which are provided by the Australian Copyright Act 1968. Users may make use of the material in the Repository providing due attribution is given and the use is in accordance with the Copyright Act 1968.

Take down policy
If you believe this document infringes copyright, raise a complaint by contacting repository-lib@uwa.edu.au. The document will be immediately withdrawn from public access while the complaint is being investigated.
Compartmentalization of innate immune responses in the central nervous system during cryptococcal meningitis/HIV coinfection. / Naranbhai, V.; Chang, C.; Durgiah, R.; Omarjee, S.; Lim, Andrew; Moosa, M.Y.S.; Elliot, J.H.; Ndung’U, T.P.; Lewin, S.R.; French, Martyn; Carr, W.H.

© 2014 Lippincott Williams & Wilkins, Inc.

This is a non-final version of an article published in final form in AIDS, Vol. 28, No. 5, p. 657-666.
The definitive published version (see citation above) is located on the journal home page of the publisher, Lippincott Williams & Wilkins.

This version was made available in the UWA Research Repository on the 13th of March 2015, in compliance with the publisher’s policies on archiving in institutional repositories.

Use of the article is subject to copyright law.
Full Title: Compartmentalisation of innate immune responses in the central nervous system during cryptococcal meningitis/ HIV co-infection

Running head: Innate immunity in Cryptococcal meningitis.

Vivek NARANBHAI1,2,3§, Christina C. CHANG2,4,5, Raveshni DURGIAH2,
Saleha OMARJEE2, Andrew LI6, Mahomed-Yunus S. MOOSA7, Julian H.
ELLIO4,5, Thumbi NDUNG’U2,8,9, Sharon R. LEWIN4,5, Martyn A. FRENCH6,
William H. CARR2,10§

Institute(s): 1Centre for the AIDS Programme of Research in South Africa, Nelson R Mandela School of Medicine, University of KwaZulu Natal, Durban, South Africa, 2HIV Pathogenesis Programme, Nelson R Mandela School of Medicine, University of KwaZulu Natal, Durban, South Africa, 3Nuffield Department of Medicine, University of Oxford, Oxford, United Kingdom, 4Department of Infectious Diseases, Monash University and Alfred Hospital, Melbourne, Australia, 5Centre for Biomedical Research, Burnet Institute, Melbourne, Australia, 6School of Pathology and Laboratory Medicine, University of Western Australia, Perth, Australia, 7Department of Infectious Diseases, Nelson R Mandela School of Medicine, University of KwaZulu Natal, Durban, South Africa, 8KwaZulu-Natal Research Institute for Tuberculosis and HIV(K-RITH), University of KwaZulu Natal, Durban, South Africa 9Max Planck Institute for Infection Biology, Berlin, Germany, 10Medgar Evers College(City University of New York), Brooklyn, United States.

Sources of Funding: This study was supported by the South African HIV/AIDS Research Platform(SHARP), the REACH initiative grant 2007 and US National Institutes for Health FIC K01-TW007793. VN was supported by LIFELab and the Columbia University-South Africa Fogarty AIDS International Training and Research Program(AITRP, grant #D43 TW000231). CCC was supported by an Australian Postgraduate Award 2009, Australian National Health and Medical Research Council(NHMRC) Postgraduate Scholarship 2010-2012. SRL is a NHMRC Practitioner Fellow. TN holds the South African Research Chair in Systems Biology of HIV/AIDS and is a Howard Hughes Medical Institute International Early Career Scientist. Additional training was supported by the South African National Research Foundation KISC Award.

Word count: Abstract: 250

Text: 3500/3500

§Corresponding authors

Vivek Naranbhai, Nuffield Department of Medicine, University of Oxford, Oxford, OX3 7BN, UK. Email: vivekn@well.ox.ac.uk & William H Carr, Department of Biology, Medgar Evers College, The City University of New York, Brooklyn, New York, 11225 USA. Email: wcarr@mec.cuny.edu
Abstract

Objective: The role of innate immunity in pathogenesis of cryptococcal meningitis (CM) is unclear. We hypothesised that NK cell and monocyte responses are central nervous system (CNS) compartmentalised, and altered by anti-fungal therapy and combination antiretroviral therapy (cART) during CM/HIV co-infection.

Design: Sub-study of a prospective cohort study of adults with CM/HIV co-infection in Durban, South Africa.

Methods: We used multi-parametric flow cytometry to study compartmentalisation of subsets, activation (CD69\textsuperscript{pos}), CXCR3 and CX3CR1 expression and cytokine secretion of NK cells and monocytes in freshly collected blood and cerebrospinal fluid (CSF) at diagnosis (n=23), completion of anti-fungal therapy induction (n=19) and after a further 4 weeks of cART (n=9).

Results: Relative to blood, CSF was enriched with CD56\textsuperscript{bright} (immunoregulatory) NK cells (p=0.0004). At enrolment, CXCR3 expression was more frequent amongst blood CD56\textsuperscript{bright} than either blood CD56\textsuperscript{dim} (p<0.0001) or CSF CD56\textsuperscript{bright} (p=0.0002) NK cells. Anti-fungal therapy diminished blood (p<0.05) but not CSF CXCR3\textsuperscript{pos} NK cell proportions nor CX3CR1\textsuperscript{pos} NK cell proportions. CD56\textsuperscript{bright} and CD56\textsuperscript{dim} NK cells were more activated in CSF than blood (p<0.0001). Anti-fungal therapy induction reduced CD56\textsuperscript{dim} NK cell activation in CSF (p=0.02). Activation of blood CD56\textsuperscript{bright} and CD56\textsuperscript{dim} NK cells was diminished following cART commencement (p<0.0001, p=0.03). Immunoregulatory NK cells in CSF tended to secrete higher levels of CXCL10 (p=0.06) and lower levels of TNF-\alpha (p=0.06) than blood immunoregulatory NK cells. CSF was enriched with non-classical monocytes (p=0.001), but anti-fungal therapy restored proportions of classical monocytes (p=0.007).

Conclusions: These results highlight CNS activation, trafficking and function of NK cells and monocytes in CM/HIV and implicate immunoregulatory NK cells and pro-inflammatory monocytes as potential modulators of CM pathogenesis during HIV co-infection.

Keywords: Cryptococcal meningitis, cerebrospinal fluid, Natural Killer cells, Monocytes, HIV-1
Cryptococcal meningitis (CM) is a major cause of morbidity and mortality in patients with HIV and AIDS. Annually, approximately 957,900 cases of CM occur, resulting in 624,700 deaths by three-months after infection, with sub-Saharan Africa bearing the largest burden of disease[1]. The underlying mechanisms causing death and disability include development of persistently high intracranial pressures, vasculopathies, and local brain inflammation with bystander neuronal damage. Both innate and adaptive immune responses contribute to the immunopathogenesis of CM but the regulation and timing of their development remain poorly understood.

Natural Killer (NK) cells are key effectors of innate immunity that are able to mediate pathogen elimination by directly killing or modulating innate and adaptive immune responses through secretion of cytokines. In humans, expression of CD56, but a lack of CD3, CD14, and CD19, defines their phenotype. Functionally, they can be further subdivided into subsets with primarily cytokine-secretion capabilities (CD56\(^{bright}\)CD16\(^{neg}\)) or cytolytic capabilities (CD56\(^{dim}\)CD16\(^{pos}\))[2]. Typically, CD56\(^{bright}\) NK cells are more prevalent at extravascular sites than CD56\(^{dim}\) NK cells[2]. During HIV disease a third subset, CD56\(^{neg}\)CD16\(^{pos}\), increases disproportionately in blood, but this subset is deficient in both cytokine production and cytolysis[3]. In vivo mouse and in vitro human studies suggest that NK cells are able to directly kill cryptococci by perforin-mediated cytotoxicity[4, 5], or indirectly by the potentiation of macrophage anti-fungal activity[6]. NK cells are able to enter the central nervous system (CNS) during inflammatory disease such as multiple sclerosis (MS)[7]; indeed they have been shown to play a major role in a variety of CNS infections[8]. Therefore, it is plausible that in vivo, NK cells may traffic to the site of cryptococcal infection and exert anti-fungal activity. Alternatively, NK cells may secrete immunoregulatory cytokines that affect recruitment and function of other innate and adaptive immune cells. The phenotype, function and mechanisms of NK cell infiltration/trafficking into the CNS are not well described in CM/HIV co-infection, and have only recently been examined in HIV mono-infection[9]. Thus delineating the profiles of NK
cells in the CSF during treated CM may allow identification of parameters that play a role in CM/HIV pathogenesis.

Monocytes/macrophages are a second innate immune leukocyte subset that plays a role in the pathogenesis of some inflammatory CNS diseases, and with which NK cells have substantial crosstalk. NK cells are required for monocyte differentiation into dendritic cells in several inflammatory disorders[10]. Conversely, monocytes/macrophages are able to activate NK cells through their secretion of pro-inflammatory cytokines, IL-12 and IL-18[11]. Monocytes can be divided into three functionally distinct subsets based on their relative expression of CD16 and CD14 (i.e., classical: CD14++CD16-; intermediate: CD14++CD16+; and non-classical: CD14+CD16++)[12]. Among these subsets the non-classical monocytes have the greatest capacity for secreting pro-inflammatory cytokines, such as tumor necrosis factor-alpha (TNF-α)[13], intermediate monocytes have superior reactive-oxygen species production and classical monocytes appear to have superior phagocytic function[12]. The role of monocytes in CM pathogenesis is unresolved; some reports suggest that monocytes may act as a ‘trojan horse’ allowing entry of intracellular cryptococci into the CNS[14]; others suggest that monocytes may mediate cryptococcal elimination[15]. Similar to other infections by intracellular pathogens disorders, in CM monocytes are likely required for pathogen elimination, but also to harbor pathogens intracellularly and impose clinically relevant immunopathology with their activity in CM[16].

Here we aimed to identify changes in the innate immune response in blood and CSF in patients with CM and HIV in South Africa. We prospectively characterised blood and CSF NK cell phenotypes, monocyte subsets and, to a lesser extent NK cell function in patients with HIV/CM co-infection at admission for care, after induction of anti-fungal therapy, and after a further 4 weeks following commencement of combination antiretroviral therapy (cART) in some patients.

**Methods**
This study was conducted as a sub-study of the Cryptococcal Immune Restoration Disease (IRD) study, which has been described previously[17]. We prospectively enrolled consenting cART-naïve, HIV-infected adults with a first-episode of microbiologically-confirmed CM at the King Edward VIII Hospital in Durban, South Africa. Briefly, whole blood and CSF were obtained at enrolment (median 2 days after diagnosis, range 0-8 days) from 23 participants. Amphotericin B was commenced immediately on diagnosis for a protocolled time of 14 days. About half of all patients had persistent cryptococcal growth after Amphotericin B therapy[17]. Following this induction period of anti-fungal therapy, 19 patients were re-sampled for blood and CSF (median 14 days after diagnosis range 10-15 days) and were commenced on cART as per contemporary guidelines[18] and continued on oral fluconazole. After 4 weeks of cART a final whole blood specimen was obtained from a subset of 9 patients based on their availability. Serial therapeutic lumbar punctures where conducted as required for therapeutic purposes whilst continuing anti-fungal therapy.

At enrolment, the mean age of participants in this sub-study was 34.7 years (range 21-55 years), and 43% were female; similar to the overall cohort[17]. The median baseline CD4+ T-cell count was 22 cells/mm³ (IQR 6.5-43), and the median plasma HIV viral load was $3.18 \text{Log}_{10}$ copies/ml (IQR 1.14-5.95). After 4 weeks of cART, amongst 9 participants from whom blood samples were available, the median CD4+ T-cell count was 74 cells/mm³ (IQR 49-153), and the median plasma HIV viral load was $2.56 \text{Log}_{10}$ copies/ml (IQR 2.31-2.91).

This study was approved by the University of KwaZulu-Natal Biomedical Research Ethics Committee, the Monash University ethics committee and the University of Western Australia ethics committee.

**Flow Cytometry analyses**

The cellular profile of CSF can be taken as a measure of cells in an intermediate compartment between blood and CNS parenchyma. We used methods that have been previously used to study CSF T-cell profiles in...
healthy and HIV-infected patients to characterise NK cells and monocytes in the CNS[19-21]. Peripheral blood and CSF leucocytes were simultaneously stained with a panel of fluorophore-conjugated antibodies and subjected to multiparametric flow cytometry using conventional methods. Briefly, for whole blood staining, 150μl undiluted whole blood collected in tubes containing ethylenediaminotetraacetic acid(EDTA) was incubated with the following antibodies for 20 minutes at 4°C: anti-CD3 APC, anti-CD8 Qdot 655, anti-CD14 Pacific Orange(PO), anti-CD16 Pacific Blue(PB), anti-CD45 AlexaFluor700(AF700), anti-CD56 PC7(Beckman Coulter, Pasadena, USA) anti-CD69 FITC, anti-CX3CR1 PE and anti-CXCR3 PerCP-Cy5.5. All antibodies were from Becton Dickinson(Franklin Lakes, USA) unless otherwise indicated. Red blood cells were lysed with VersaLyse( Beckman Coulter) as per the manufacturer’s directions, pelleted by centrifugation and fixed with a paraformaldehyde-containing fixative(Reagent A, Life Technologies, Carlsbad, USA). For CSF cell staining, the total volume of CSF obtained from the patient(ranging from 3-30 ml), was centrifuged at 750 x g for 10 minutes, resuspended in 1ml R10(RPMI 1640 containing penicillin/streptomycin, 10% fetal calf serum, supplemented with 1.0 mg/ml L-glutamine) and live nucleated cells were enumerated with a nucleocounter(Chemometec, Allerod, Denmark). The range of the nucleocounter was 5x10^3-2x10^6 cells/ml and since several samples were outside this range we were unable to convert proportions to absolute numbers in this study. One third of the total number of nucleated cells were aliquoted into FACSTubes, washed with Dulbeco’s phosphate buffered saline(DPBS) and stained for 20 minutes with the same panel of antibodies listed above. Cells were washed and fixed as above but the lysis step was omitted.

For intracellular cytokine staining experiments: the primary stain included anti-CD3 APC( Beckman Coulter), anti-CD56 PC7( Beckman Coulter) and anti-CD16 PB. Following fixation, cells were incubated for 15 minutes, washed, and then permeabilised and stained with anti-cytokine antibodies by adding Reagent B(Life Technologies), anti-CXCL10 PE and anti-TNFα AF700. After a further 15 minutes cells were washed with DPBS.

Flow cytometry data were collected on a BD LSRII and analyzed using FlowJo v10.0.6(Treestar, Ashland, USA). At least 5,000 CD45+ leucocytes
were collected for each CSF sample, and $3 \times 10^6$ events were collected for each whole blood specimen. Fluorescence minus one gating strategies were used to determine gating boundaries as described[22]. The gating strategy is shown in Supplementary Figure 1.

**Statistical analyses**

For comparisons between paired specimens, from the same individual at different time-points, or at the same time-point but from blood and CSF, a non-parametric matched-pairs Wilcoxon signed rank test was performed. This method ignores data points where the pair is incomplete and thus is robust to missing data for the 4 individuals for whom samples were unavailable at completion of anti-fungal therapy induction. Statistical analyses were conducted in GraphPad Prism v5(GraphPad, La Jolla, California).

**Results**

**Proportions of immunoregulatory NK cells(CD56\text{bright}) are expanded in the CSF of patients with CM and HIV**

Cytolytic and cytokine-secretory roles are performed by different NK cell subsets that partially overlap in function: low expression of CD56(CD56\text{dim}) demarcates cytolytic NK cells and high expression of CD56(CD56\text{bright}) identifies a cytokine-secreting subset that is thought to be less mature[23]. At both enrolment and after completion of anti-fungal therapy, the CSF was enriched with CD56\text{bright} immunoregulatory NK cells compared to blood(at enrolment median 20% vs. 5.4%, median change $\Delta=13.32\%$, $p=0.0004$, Figure 1A), and had fewer CD56\text{dim} NK cells(at enrolment median 64% vs. 86.1%, median $\Delta =20.6\%$, $p<0.0001$). The ratio of CD56\text{bright} / CD56\text{dim} NK cells was significantly higher in the CSF compared to blood(Figure 1B). Neither the absolute proportions nor the ratio of CD56\text{bright} and CD56\text{dim} NK cells was significantly modified following 14 days of anti-fungal therapy(Figure 1B).
Expansion of an anergic subset of NK cells with low/absent CD56 expression (CD56$^{\text{neg}}$) has been observed in the blood of patients with advanced HIV disease[24]. Notwithstanding the use of classical methods as opposed more recently described methods that enhance specificity of NK cell gating[25], we did not observe differences in the proportion of CD56$^{\text{neg}}$ NK cells in the CSF relative to blood (Supplementary Figure 2). In both blood and CSF the median proportion of CD56$^{\text{neg}}$ NK cells was 6-7%.

In both blood and CSF, immunoregulatory (CD56$^{\text{bright}}$) NK cells and cytolytic (CD56$^{\text{dim}}$) NK cells differed in their expression of CXCR3.

To investigate whether immunoregulatory NK cells differed from cytotoxic NK cells in CSF, we compared their expression of chemokine receptors. Eisenhardt and colleagues recently demonstrated that CXCR3 expression in extravascular tissues demarcated specific NK cell subsets that play a role in infection[26]. Moreover, CXCR3 is the receptor for pro-inflammatory chemokines: CXCL9(MIG), CXCL-10(IP-10) and CXCL-11(ITAC)[27]. Based on our prior discovery of an increasing gradient of CXCL-10 from blood to CSF in the participants of this study[28], we speculated that this chemokine could mediate chemotaxis of CXCR3-expressing NK cells into the CNS.

At enrolment in blood we found a greater proportion of CD56$^{\text{bright}}$ NK cells expressing CXCR3 than CD56$^{\text{dim}}$ NK cells (median 5.4% vs. 2.2%, median $\Delta$=3.2%, $p<0.0001$; Figure 2A). In contrast, in CSF a significantly greater proportion of CD56$^{\text{dim}}$ NK cells expressed CXCR3 than CD56$^{\text{bright}}$ NK cells (median 4.2% vs. 2.4%, median $\Delta$=1.5%, $p=0.0011$). Furthermore, we found that differential CXCR3 expression on CD56$^{\text{bright}}$ and CD56$^{\text{dim}}$ NK cells extended to comparisons between blood and CSF compartments (Figures 2B and 2C). Amongst CD56$^{\text{bright}}$ NK cells, the proportion expressing CXCR3 was significantly greater in blood than CSF at enrolment (median 7.8% vs. 2.1%, median $\Delta$=4.6%, $p=0.0002$), but after 14 days of anti-fungal therapy the proportion of CXCR3$^{\text{pos}}$ CD56$^{\text{bright}}$ NK cells in blood declined (median 7.8% vs. 4.8%, median $\Delta$=2.8%, $p=0.009$). By completion of anti-fungal therapy
induction there was no difference between blood and CSF in the proportion of
CD56\textsuperscript{bright} NK cells expressing CXCR3(Figure 2B). The proportion of
CXCR3\textsuperscript{pos} CD56\textsuperscript{bright} NK cells in CSF did not change over the period of anti-
fungal therapy induction.

In contrast, amongst CD56\textsuperscript{dim} NK cells, the proportion expressing
CXCR3 was significantly greater in CSF than blood at both enrolment and
after completion of anti-fungal therapy(median 8.6% vs. 3.4%, median
\(\Delta=3.2\)% at enrolment, \(p=0.001\); median 7.9% vs. 2.5%, median \(\Delta=5.36\)% at
completion of anti-fungal therapy, \(p=0.0005\); Figure 2C). This difference in
CXCR3 expression was maintained in CSF over the period of anti-fungal
therapy induction. However, in blood the proportion of CD56\textsuperscript{dim} NK cells
expressing CXCR3 declined(median 3.4% at enrolment vs. 2.3% at
completion of anti-fungal therapy, median \(\Delta=1.2\)%, \(p=0.03\)).

The proportion of CXCR3\textsuperscript{pos} CD56\textsuperscript{bright} and CD56\textsuperscript{dim} NK cells in blood
following 4 weeks of combined antiretroviral therapy(cART) did not differ from
that at completion of anti-fungal therapy(Supplementary Figure 3A and B).

We also examined expression of CX3CR1 on the various NK cell
subsets, as CX3CR1 expressing NK cells have been reported to be involved
in modifying autoimmune CNS disease pathogenesis[29]. At enrolment, in
CSF, there was a higher proportion of CD56\textsuperscript{dim} NK cells expressing CX3CR1
than CD56\textsuperscript{bright} NK cells(median 10.6% vs. 2.9%, median \(\Delta=4.8\)%, \(p=0.0003\))
but there was no difference observed in blood. Both at enrolment and at
completion of anti-fungal therapy induction, a larger proportion of
CD56\textsuperscript{bright} NK cells in blood expressed CX3CR1 than those in CSF(median
15.2% vs. 2.8%, median \(\Delta=14.3\)%, \(p=0.001\); and median 10.4% vs. 1.4%,
median \(\Delta=6.5\)%, \(p=0.009\), respectively, Figure 2B). The proportion of
CX3CR1 expressing CD56\textsuperscript{dim} NK cells did not differ between blood and CSF
regardless of timepoint or anti-fungal therapy. In summary, these data
demonstrate that NK cells differ in expression of CXCR3 and to a lesser
extent CX3CR1 chemokine receptors according to compartment and subset.
Cytotoxic and immunoregulatory NK cells in CSF were more activated than NK cells in blood

Activation is a necessary precursor of both CD56<sup>dim</sup> and CD56<sup>bright</sup> NK cell activity. To gain insight into the role of these NK cells in CM pathogenesis we examined the proportion of activated cells in each subset by measuring cell-surface expression of CD69, an early marker of lymphocyte activation. CD56<sup>bright</sup> and CD56<sup>dim</sup> NK cells in CSF were more activated than blood NK cells (Figure 3) at enrolment (median 48.1% vs. 14.3%, median Δ=37.7%, p<0.0001; and median 54.2% vs. 19.7%, median Δ=33.3%, p<0.0001 respectively) and after completion of anti-fungal therapy induction (median 52.9% vs. 7.9%, median Δ=43.2%, p=0.0001; and median 46.5% vs. 9.95, median Δ=31.6%, p=0.0003 respectively). Although anti-fungal therapy did not significantly reduce the proportion of activated NK cells in blood, or the proportion of activated CD56<sup>bright</sup> NK cells in CSF, after completion of anti-fungal therapy induction the proportion of activated CD56<sup>dim</sup> NK cells in CSF was significantly reduced (Figure 3). After completion of anti-fungal therapy induction, the proportion of activated blood NK cells was positively associated with the plasma HIV VL (r=0.65, p=0.007). Consistent with previous reports[30, 31] cART commencement was associated with a significant decline in the proportions of CD69<sup>pos</sup> NK cells in both CD56<sup>bright</sup> median 16.1% vs. 5.7%, p<0.0001) and CD56<sup>dim</sup> <span class="red-highlight" style="color: red;">blood</span> NK-cell subsets (median 10.3% vs. 6.7%, p=0.03, Supplementary Figure 2C and 2D).

Immunoregulatory NK cells in CSF expressed higher levels of CXCL10 and lower levels of TNF-α than NK cells in blood prior to commencing anti-fungal therapy

Next, we examined whether the chemokine and cytokine secretion profiles of CD56<sup>bright</sup> NK cells in CSF differed from those in blood. CSF levels
of the chemokine CXCL-10(IP-10) levels in CSF correlate directly with
neuronal injury in CNS HIV disease[32]. Similarly, the amounts of pro-
inflammatory chemokines and cytokines, including CXCL-10 and tumor
necrosis factor-alpha(TNF-α), correlate with clinical outcomes in CM and HIV
coinfection prior to starting cART[33]. Thus, to quantify differences in
functional responses during HIV and CM co-infection we compared
intracellular cytokine profiles of CXCL-10 and TNF-α in NK cells. We obtained
paired blood and CSF samples at enrolment from five participants and
performed intracellular cytokine staining for CXCL-10 and TNF-α. The
proportion of CD56<sup>bright</sup> NK cells in CSF expressing CXCL-10 tended to be
higher than in blood(median 57.6% vs. 35.7% median Δ=19.8%, p=0.06,
Figure 4). Conversely, the proportion of CD56<sup>bright</sup> NK cells in CSF expressing
TNF-α tended to be lower than in blood(median 64.2 vs. 42.3%, median
Δ=19.84%, p=0.06, Figure 4).

**CSF was enriched for non-classical monocytes in CM prior to anti-fungal
therapy**

NK cells engage in a bi-directional communication with other innate
and adaptive immune cells. During neuroinflammation, monocytes are a major
cell type that is recruited to the CNS[34], unlike other tissues that recruit
neutrophils. Therefore we also evaluated our flow cytometric data to quantify
monocyte subsets in CSF and blood.

Relative to blood, CSF was enriched for non-classical ‘pro-
inflammatory’ monocytes at enrolment(median 3.12% vs. 0.78%, median
Δ=1.83%, p=0.001, Figure 5A). Correspondingly, the proportion of classical
monocytes was lower in CSF than blood(median 30% vs. 64%, median
Δ=59.8%, p=0.0007). There was also a trend towards a greater proportion of
intermediate monocytes in CSF than blood(median 18.1% vs. 13.5%, median
Δ=6.5%, p=0.07).
Anti-fungal therapy restores proportions of classical monocytes in CSF

Comparing the proportions of the three major monocytes subsets in CSF at enrolment and after completion of anti-fungal therapy demonstrated that the proportion of classical monocytes significantly increased over time (median 36.1% to 89.5%, median $\Delta=23.4\%$, $p=0.006$, Figure 5B). In contrast, the proportion of intermediate monocytes significantly decreased (median 13% vs. 2.69%, median $\Delta=8.4\%$, $p=0.003$), while the proportion of non-classical monocytes also declined but the difference did not achieve statistical significance (median 2.78% vs. 3.85%, median $\Delta=2.5\%$, $p=0.06$). In comparison, there were no significant differences between proportions of monocyte subsets in blood at enrolment and after 14 days of anti-fungal therapy (data not shown).

Discussion

Here we assessed NK cells and monocytes in CSF and blood in patients with HIV-CM prior to and following anti-fungal therapy induction and cART. We found that markers of activation and/or function expressed by NK cells and monocytes were compartmentalised in the CNS relative to blood. These findings suggest that immunoregulatory NK cells and non-classical monocytes may play a role in CM pathogenesis. Such changes might contribute to adverse outcomes after commencing cART, such as cryptococcosis-associated immune reconstitution inflammatory syndrome (C-IRIS).

Consistent with previous reports of the phenotype of NK cells in extravascular tissues [2] and during CNS infections [35], we found a higher proportion of immunoregulatory (CD56$^{bright}$) NK cells in CSF than in blood. Homing of plasmablasts and T-cells to the CNS compartment have been shown to be mediated by CXCR3 [36, 37], a receptor for CXCL10. We previously reported that there was a higher concentration of CXCL-10 in CSF than in blood in this cohort [28]. CX3CL1 has also been reported to mediate migration of NK cells to the CNS during experimental allergic
encephalomyelitis[29]. We observed differences between blood and CSF in the proportions of both CXCR3\textsuperscript{pos} and CX3CR1\textsuperscript{pos} NK cells. We therefore speculate that the enhanced CXCR3 and/or CX3CR1 expression on CD56\textsuperscript{bright} NK cells in blood that we observed may have equipped these cells to enter the CNS compartment in response to a CXCL-10 or CX3CL1 gradient. Tracking chemokine expression on particular cells to test this hypothesis was beyond the scope of our work, and remains to be tested.

It is notable that the proportion of CXCR3 expressing NK cells in the CSF was not affected by antifungal therapy induction. This subset has been reported to have impaired cytotoxic and cytokine-secretory capacity in Hepatitis C virus infection[26]. The maintenance of this population of NK cells in CSF may have implications in the development of adverse clinical sequelae such as C-IRIS. Understanding of role of this subset in CM infection will benefit from further study.

We extended previous reports of NK cells in the CNS by assessing their cytokine profiles and activation status over time. As predicted, our findings were consistent with previous reports of decreased activation of NK cells in blood of HIV-infected adults following cART initiation and the reduction of HIV burden[30, 31]. However, we observed a partial decline in NK cell activation only in the CD56\textsuperscript{dim} subset in the CNS following two weeks of antifungal therapy. We attributed the maintenance of NK cell activation in the CNS to either residual pathogen burden in the CNS [17] or an intrinsic high threshold for deactivation.

Our preliminary discovery that immunoregulatory NK cells secreted more CXCL-10 in CSF than in blood, suggested that they were preferentially promoting a pro-inflammatory environment in the CNS compartment. The observation that the immunoregulatory NK cells in CSF produced less TNF than in blood may be a result of interaction with dendritic cells or with cryptococci[38]. Taking into consideration that NK cells also generate strong IFN-\(\gamma\) responses to \textit{C. neoformans}[5] and that IFN-\(\gamma\) levels in CSF are one of the leading predictors of clinical outcomes in CM pathogenesis[33], they could be candidates for immune modulation to improve clinical outcomes in this
disease. However, we examined only a small number of patients and further studies are needed.

In contrast to NK cells, we observed a rebalancing of monocyte subsets following anti-fungal therapy induction. After 14 days of anti-fungal therapy the proportions of intermediate monocyte subsets in CSF declined, whereas the proportion of classical monocytes increased. We attribute this change to differences in functional roles during clearance of Cryptococcus. Unlike classical monocytes, non-classical and intermediate monocytes secrete large amounts of TNF-α and IL-1β, are expanded during many infectious diseases[39-41] including HIV[42], and are preferentially recruited to sites of inflammation[16]. Restoration of monocyte subset distribution with anti-fungal therapy suggests that reducing antigen burden is sufficient to restore monocyte homeostasis in the CNS compartment.

Although we discovered novel changes in NK cells and monocytes phenotypes in the CNS compartment during treated CM disease, our findings have limitations. Because we were unable to quantify absolute numbers of cells in either CSF or in blood, we cannot infer whether absolute numbers of specific subsets were altered. With the exception of anti-fungal therapy induction, we were unable to examine the association between innate immunological events in CSF or blood and clinical outcomes. Nor can we definitely demonstrate whether these observations are specific to HIV/CM or may be observed in other forms of meningoencephalitis with or without HIV infection. We examined the effect of cART in only 9 patients. Larger studies are required to establish the clinical relevance of our findings. Nevertheless, our data provided new insights into regulation of compartmentalised immune responses during treated CM disease in adults with advanced HIV.

In summary, our findings suggest that NK cell and monocyte responses to cryptococci are compartmentalised in patients with CM and HIV co-infection. Furthermore, they highlight a potential role of immunoregulatory NK cells and different monocyte subsets in CM pathogenesis. Prospective studies of CNS-resident NK cells and monocytes, and their association with clinical outcomes, such as C-IRIS, are warranted.
Acknowledgements

We would like to thank the staff of the HIV Pathogenesis Programme (HPP) at the University of KwaZulu-Natal (Durban, South Africa) for their assistance in processing clinical samples.

Author Contributions

VN and CC conceived and conducted the experiments described in this study. RD, SO, and AL provided technical support for experiments. MYS, JHE, TN, SR, MAF and WHC provided overall oversight of the clinical cohort accrual, follow-up and experimental procedures. MAF and WHC provided critical advice throughout the conduct of the study. All the authors read, commented and approved the final version of this manuscript.
Figure 1. At enrolment and after completion of anti-fungal therapy induction, CSF was enriched with CD56\textsuperscript{bright} NK cells, had fewer CD56\textsuperscript{dim} and similar frequencies of CD56\textsuperscript{neg} NK cells relative to blood (a) and hence the ratio of %CD56\textsuperscript{bright} (immunoregulatory) NK cells to %CD56\textsuperscript{dim} (cytotoxic) NK cells was higher in CSF than blood at enrolment and after completion of anti-fungal therapy induction in HIV patients with CM(a). Medians (horizontal lines) and interquartile ranges (whiskers) are shown in each graph. Measurements in blood denoted by black squares (■) and in CSF denoted by grey circles (●).

Figure 2. CXCR3 expression differed by NK cell subset (CD56\textsuperscript{bright}, CD56\textsuperscript{dim}) and compartment. In blood, at enrolment the proportion of NK cells expressing CXCR3 was higher among CD56\textsuperscript{bright} NK cells than among CD56\textsuperscript{dim} NK cells, but the opposite was observed in CSF(a). The proportion of CD56\textsuperscript{bright} NK cells expressing CXCR3 was significantly higher in blood at enrolment but declined following anti-fungal therapy. By completion of anti-fungal therapy the proportion in blood was similar to CSF(b). In contrast, the proportion of CD56\textsuperscript{dim} NK cells expressing CXCR3 was significantly higher in CSF than blood at enrolment and completion of anti-fungal therapy(c). Medians (horizontal lines) and interquartile ranges (whiskers) are shown in each graph. Measurements in blood denoted by black squares (■) and in CSF denoted by grey circles (●).
Figure 3. CD56\textsuperscript{bright}(a) and CD56\textsuperscript{dim}(b) NK cells were more activated in CSF than in blood at enrolment and completion of anti-fungal therapy, and activation was only partially reduced by anti-fungal therapy induction. Medians (horizontal lines) and interquartile ranges (whiskers) are shown in each graph. Measurements in blood denoted by black squares (■) and in CSF denoted by grey circles(●).

Figure 4. At enrolment the proportion of CD56\textsuperscript{bright} NK cells producing CXCL-10 was higher in CSF than in blood, but the proportion producing TNF-\(\alpha\) was lower in CSF than in blood\((n=5)\). Results are expressed as the percentage of cytokine producing cells. Measurements in blood denoted by black squares (■) and in CSF denoted by grey circles(●).

Figure 5. The proportions of non-classical and intermediate monocytes in CSF declined following anti-fungal therapy induction, whereas the proportion of classical monocytes increased. At enrolment CSF was enriched with non-classical monocytes\((CD14^{lo}CD16^{hi})\) and intermediate monocytes\((CD14^{hi}CD16^{lo})\), but had significantly lower proportions of classical monocytes\((CD14^{hi}CD16^{neg})\)(a). After completion of anti-fungal therapy induction the proportions of classical monocyte in CSF were increased and the proportions of intermediate and non-classical monocytes in CSF were reduced(b). Results are expressed as the percentage of total monocytes. Medians (horizontal lines) and interquartile ranges (whiskers) are shown in each graph. Measurements in blood denoted by black squares (■) and in CSF denoted by grey circles(●).
Supplementary Figure 1: Gating strategy for Natural Killer (NK) cells (a) and monocytes (b). NK cells were gated as singlets, CD45^{pos}, CD14^{neg}, FSC-SSC appropriate for lymphocytes, CD3^{neg}, CD16/CD56^{pos} cells and then gated further as CD56^{bright}, CD56^{dim} or CD56^{neg} NK cells. Monocytes were similarly gated as singlet, CD45^{intermediate}/SSC appropriate for monocytes, CD3^{neg}, CD14/CD16^{pos} cells and then gated further as classical (CD14^{hi}/CD16^{neg}), intermediate (CD14^{hi}/CD16^{lo}) or non-classical (CD14^{lo}/CD16^{hi}) monocytes. Gating of CXCR3, CX3CR1 or CD69 expression was performed using fluorescence-minus-one gates (FMO) to set lower thresholds for gating.

Supplementary Figure 2: Four weeks of cART therapy did not affect CXCR3 expression on CD56^{bright} (a) or CD56^{dim} (b) NK cells, but reduced the proportions of CD69^{pos} activated NK cells in both CD56^{bright} (c) and CD56^{dim} (d) NK cell fractions. Week 0 refers to start of cART and Week 4 refers to sampling after 4 weeks of cART. Results are expressed as the percentage of total monocytes. Medians (horizontal lines) and interquartile ranges (whiskers) are shown in each graph. Measurements in blood denoted by black squares (■) and in CSF denoted by grey circles (○).
Supplementary Figure 3: CX3CR1 expression differed by NK cell subset (CD56<bright, CD56<sub>dim</sub>) and compartment. In blood, at enrolment the proportion of NK cells expressing CX3CR1 was similar between CD56<bright and CD56<sub>dim</sub> NK cells, but in CSF, a larger proportion of CD56<bright NK cells expressed CX3CR1 compared with CD56<sub>dim</sub> NK cells(a). The proportion of CD56<bright NK cells expressing CX3CR1 was significantly higher in blood at enrolment and after completion of anti-fungal therapy induction(b). Amongst CD56<sub>dim</sub> NK cells, no differences in CX3CR1 expression between CSF and blood were noted at enrolment or after completion of anti-fungal therapy.(c). Medians (horizontal lines) and interquartile ranges (whiskers) are shown in each graph.
Figure 1

A

Enrolment

Antifungal therapy induction period

Completion

B

Enrolment

Antifungal therapy induction period
Figure 2

A

\[
\begin{align*}
\text{%CXCR3\textsuperscript{+}} & \text{NK cells} \\
\text{CD56\textsuperscript{dim}} & \quad \text{CD56\textsuperscript{bright}} \\
\text{Blood} & \quad \text{CSF}
\end{align*}
\]

\(p < 0.0001\) \quad \(p = 0.0011\)

B

\[
\begin{align*}
\text{%CXCR3\textsuperscript{+}} & \text{CD56\textsuperscript{bright} NK cells} \\
\text{Blood} & \quad \text{CSF} \\
\text{Enrolment} & \quad \text{Completion}
\end{align*}
\]

\(p = 0.0002\) \quad \(p = 0.0092\)

C

\[
\begin{align*}
\text{%CXCR3\textsuperscript{+}} & \text{CD56\textsuperscript{dim} NK cells} \\
\text{Blood} & \quad \text{CSF} \\
\text{Enrolment} & \quad \text{Completion}
\end{align*}
\]

\(p = 0.0012\) \quad \(p = 0.0005\)

Antifungal therapy induction period
Figure 3

A

B

Antifungal therapy induction period
Figure 5.

A

% Monocytes

Blood Classical | Blood Intermediate | Blood Non-classical

CSF

p = 0.0007  p = 0.0731  p = 0.0011

B

% Monocytes

Enrolment Completion Classical | Enrolment Completion Intermediate | Enrolment Completion Non-classical

p = 0.0067  p = 0.0034  p = 0.0580
Supplementary Figure 1

A

B

<Pacific Blue-A>: CD16

<PerCP-Cy5.5-A>: CXCR3

<FITC-A>: CD69

<APC-A>: CD3

<APC-A>: CD3

<APC-A>: CD3

<APC-A>: CD3

<APC-A>: CD3

<APC-A>: CD3
Supplementary Figure 3

A

![](image)

B

![](image)

C

![](image)
References


30


37. Stiles LN, Hosking MP, Edwards RA, Strieter RM, Lane TE. Differential roles for CXCR3 in CD4+ and CD8+ T cell trafficking following viral infection of the CNS. *Eur J Immunol* 2006, **36**:613-622.


