Measuring Soluble ICAM-1 in African Populations

Abdirahman I. Abdi1,3*, Michelle Muthui1, Esther Kiragu1, Peter C. Bull1,2

1 KEMRI-Wellcome Trust Research Programme, Kilifi, Kenya, 2 Nuffield Department of Clinical Medicine, John Radcliffe Hospital, University of Oxford, Oxford, United Kingdom, 3 Department of Biochemistry and Chemistry, Pwani University, Kilifi, Kenya

Abstract

The level of plasma soluble ICAM-1 (sICAM-1) has been associated with the pathogenesis of several diseases. Previously, a commercial antibody was reported not to recognize an ICAM-1 allele known as ICAM-1kilifi prevalent among African populations. However, that study was based on 19 samples from African Americans of whom 13 had the wild type allele, five heterozygotes and one homozygote. Here, we compare plasma sICAM-1 measures using three different commercial antibodies in samples from Kenyan children genotyped for ICAM-1kilifi allele. We show that two of these antibodies have some degree of deficiency in detecting the ICAM-1kilifi allele. Consideration of the antibody used to measure sICAM-1 is important as up to 30% of the populations in Africa harbour this allele.

Introduction

Adhesion molecules expressed on vascular endothelial cells in response to inflammation are crucial for the pathogenesis of infectious (reviewed in [1]) and non-infectious diseases [2]. One well studied molecule is the intercellular adhesion molecule-1 (ICAM-1) expressed on endothelial cells where its expression is amplified by pro-inflammatory cytokines [3]. ICAM-1 plays a role in recruitment of leukocytes to the vascular wall in response to inflammation [4]. In malaria, it plays a role in the sequestration of infected erythrocytes in the organs [5], leading to blockage of blood vessels and coma. ICAM-1 is also a receptor for Rhinovirus [6].

ICAM-1 protein is found in both membrane-bound and soluble forms. The soluble form is thought to arise from active cleavage of the endothelial membrane-bound form by a zine dependent metalloprotease [7]. Soluble levels of ICAM-1 (sICAM-1) in plasma have been associated with coronary heart disease and other vascular diseases [2]. In malaria, sepsis, and other infectious diseases, increasing plasma sICAM-1 has been associated with severity of disease (reviewed in [1]).

Polymorphisms within the ICAM1 coding gene have been shown to influence the level of plasma sICAM-1 [8]. Of relevance to this report is a single nucleotide point mutation (SNP) at the rs5491 first reported by Fernandez-Reyes et al [9], who sequenced the coding region from Kenyan children. This SNP leads to a non-synonymous mutation that brings about a change from lysine to methionine (K29M) and this variant is referred to as ICAM-1kilifi. The frequency of this allele is around 20–30% in many African populations [9–12] and is rare among Caucasian populations [13].

A study by Register et al [13] first reported that a monoclonal antibody from R&D Systems (BBE1B) does not recognize ICAM-1kilifi. Another study using the same R&D Systems antibody found that the levels of sICAM-1 reduces by 50% in the heterozygotes (K29/M29), while the homozygotes (M29/M29) have levels below detection [8]. To examine whether the low sICAM-1 levels observed with the R&D monoclonal antibody is due to inability of the antibody to recognize the ICAM-1kilifi allele rather than unavailability of the protein, Register et al [13] used an alternative monoclonal sICAM-1 (BMS201INST; Bender MedSystems) in ELISA. With this alternative antibody, sICAM-1 was recognized to a greater extent in plasma samples from both heterozygotes and homozygotes suggesting that the almost background sICAM-1 levels observed in the homozygotes (M29/M29) with the R&D antibody (BBE1B) is likely due to its inability to detect the ICAM-1kilifi allele. However, the sample size in Register’s study was small where the homozygotes were represented by only one sample and therefore we sought to confirm this finding in a larger sample size.

It was also reported, using another antibody from R&D systems (cyto, DY720), that the plasma levels of sICAM-1 are not affected by the ICAM-1kilifi allele [14], though no data was given. Since the original R&D antibody BBE1B showing marked variation between alleles is no longer marketed under that name, we performed a comparative study of plasma sICAM-1 measures using two antibodies from R&D (DCD540 and DY720) and the antibody from Bender MedSystems (BMS201MST) in a group of Kenyan children genotyped for ICAM-1kilifi allele. We show that the antibody in the kit DCD540 is hardly able to detect the ICAM-1kilifi allele whilst the BMS201MST antibody has reduced ability to detect this allele. We confirm that DY720 antibody is able to detect sICAM-1 equally for both alleles.

Methods

Ethics statement

Ethical approval was obtained from Kenya Medical Research Institute (KEMRI) Ethical Review Committee (SSC 1131), and
written informed consent was obtained from parents/Guardians of the study participants.

ICAM-1 genotyping and sicam1 ELISA

Samples used in this study are from children presenting with malaria to Kilifi county hospital, a rural town located at the coast of Kenya. The children were recruited between August 2003 and September 2007 [15]. We used published methods for host genotyping of the K29M polymorphism of ICAM1 [16] and primers described in [10].

Two ELISA kits from R&D were used; catalog numbers DCD540 and DY720 and one from Bender MedSystems; catalog

Figure 1. Plasma sICAM-1 in relation to ICAM-1**Kilifi** genotypes. A. Venn diagram showing the samples used in this study. We measured sICAM-1 in 163, 150, and 58 samples using DY720, BMS201MST, and DCD540 ELISA kits. These samples were genotyped from ICAM-1**Kilifi** mutation. Of these, 145 samples have been measured with both BMS201MST and R&D DY720, 52 with R&D DY720 and R&D DCD540, 48 with BMS201MST and R&D DCD540, and 45 with all the three kits. B. Plasma sICAM1 obtained with three different kits relative to ICAM-1**Kilifi** genotypes. C. Plasma sICAM-1 in a subset of 45 samples where a measure was obtained with each of the three antibodies used. Measures of plasma sICAM-1 obtained with the BMS201MST, R&D (DCD540) and R&D (DY720) antibodies are depicted in black, red and green triangles respectively. The horizontal bar indicates the median sICAM-1 value. P value was calculated using Cuzick’s test for trend. doi:10.1371/journal.pone.0108956.g001
Table 1. Published work done in Africa that measure sICAM-1.

| Article | Article Kit used to measure sICAM-1 |
|---------|-------------------------------------|
| Bellamy R et al, Trans R Soc Trop Med Hyg 1998 [10] | R&D DY720 |
| Tchinda VH et al Acta Trop 2007 [18] | R&D DY720 |
| Conroy AL et al Plos One 2010 [19] | R&D DY720 |
| Csereti-Gazdewich CM et al, Malar J. 2010 [20] | R&D DY720 |
| Adulko, S et al, Plos One 2013 [21] | R&D DY720 |
| Djoba Siwaya JJ et al, J infect 2008 [22] | R&D but not specified |
| Park GS et al, J Pediatric Infect Dis Soc 2012 [23] | R&D but not specified |
| Mita-Mendoza, NK et al Plos One 2013 [25] | R&D LAD000 (Human Adhesion Molecule MultiAnalyte Profiling Base Kit) |
| El-Deek SE et al Med Princ Pract 2013 [26] | Bender MedSystem GmbH, Campus Vienna Biocenter |

doi:10.1371/journal.pone.0108956.t001

Results and Discussion

Plasma sICAM-1 for 58, 163 and 150 samples genotyped for ICAM-1 K29M polymorphism were obtained using the DCD540, DY720 and BMS201MST ELISA kits respectively as shown in figure 1A. Of these, 45 samples had a measure of sICAM-1 with each of the three kits (Fig. 1A).

Initially, we used the DCD540 antibody to measure sICAM-1 in 58 plasma samples genotyped for ICAM-1\textsuperscript{kilifi}. We found that the plasma sICAM-1 levels decrease from wild-type ICAM-1\textsuperscript{kilifi} heterozygotes and homozygotes in that order (Figure 1B). This decreasing trend in the levels of sICAM-1 was statistically significant (P<0.0001, Cuzick’s test for trend). This result is similar to that previously reported with the R&D (BBE1B) monoclonal antibody [13].

The Register et al [13] study used another Kit (BMS201INST) to demonstrate that expression of sICAM1 was not altered by the ICAM-1\textsuperscript{kilifi} allele [13]. However this was difficult to interpret, as the sample size was small. We therefore measured sICAM-1 in 150 samples genotyped for ICAM-1\textsuperscript{kilifi} allele using the BMS201MST antibody. With this antibody, although detection of sICAM-1 improved in both heterozygotes and homozygotes, we still observed a significant decreasing trend in the levels when moving from 0 to 1 to 2 copies of the ICAM-1\textsuperscript{kilifi} allele (Fig. 1B; P = 0.04, Cuzick’s test for trend) suggesting a deficiency in the ability of BMS201MST to detect ICAM-1\textsuperscript{kilifi} allele.

Finally, as a second alternative to BMS201MST we measured plasma sICAM-1 in 163 samples genotyped for ICAM-1\textsuperscript{kilifi} allele using the DY720 antibody from R&D Systems. With this antibody, the median levels were the same in the different ICAM-1\textsuperscript{kilifi} genotypes (Fig. 1B) confirming that the antibody in this kit is not affected by the presence of the ICAM-1\textsuperscript{kilifi} allele [14]. To directly compare the BMS201MST, and DY720, we compared their performance in the 145 samples with sICAM-1 measures with both antibodies. Again, while the ICAM-1 levels obtained with BMS201MST showed a significant decreasing trend from wild type to heterozygotes to homozygotes (N = 145, P = 0.03, Cuzick’s test for trend), the measures obtained with DY720 showed no difference (N = 145, P = 1, Cuzick’s test for trend). A similar trend was observed with 45 samples for which we had a measure of plasma sICAM-1 against all the three antibodies (Fig. 1C).

The result obtained with DCD540 antibody used in this study is largely consistent with that described in Register et al [13] using the BBE1B monoclonal antibody. Our result also suggests that the monoclonal antibody in the BMS201MST kit has some deficiency in detecting ICAM-1\textsuperscript{kilifi} allele (Fig. 1B). Of the three antibodies we tested, the DY720 antibody of the R&D Systems seems to be the ideal for measuring sICAM-1 levels in samples from African individuals and the DCD540 to be the least effective.

As the frequency of ICAM-1\textsuperscript{kilifi} allele is reported to be as high as 30% in some parts of Africa, we sought to know the number of studies carried in Africa that measured sICAM-1 and what kind of antibodies they used (summarised in table 1). We found that the majority of the studies were done in the context of malaria and used the DY720 antibody [18–21]. Other studies stated that they used a kit from R&D without specifying [22,23].

Since, in a large study, the ICAM-1\textsuperscript{kilifi} polymorphism was shown not to be associated with severe malaria [12], it is unlikely that this would have altered the conclusions in studies with severe malaria (and this is supported by a personal communication from Parks GS & John CC). Nevertheless, the high prevalence of this mutation among African populations raises the possibility that it is selected for by other conditions prevalent in Africa including other infectious diseases. This is supported by a previous study that found an association between ICAM-1\textsuperscript{kilifi} allele and reduced incidence of non-malarial febrile illness in Kenyan children [24]. Therefore, consideration of the effect of ICAM-1\textsuperscript{kilifi} allele is important because of its impact on the performance of some of the commercial ELISA kits used to measure sICAM-1.

Acknowledgments

We are grateful to the children and their parents/guardians for participation in the study. We thank Dr Gregory Fegan for his statistical advice. This work was supported by Wellcome Trust Programme Grants (084535 and 077092 to PCB and Prof Kevin Marsh (KM)) and Project...
Author Contributions
Conceived and designed the experiments: AA PB. Performed the experiments: MM EK. Analyzed the data: AA PB. Wrote the paper: AA PB.

References
1. Page AV, Liles WC (2013) Biomarkers of endothelial activation/dysfunction in infectious diseases. Virulence 4: 507–516.
2. Ridker PM, Hennekens CH, Reitman-Johnson B, Stampfer MJ, Allen J (1998) Plasma concentration of soluble intercellular adhesion molecule 1 and risks of future myocardial infarction in apparently healthy men. Lancet 351: 80–92.
3. Dietrich JB (2002) The adhesion molecule ICAM-1 and its regulation in relation with the blood-brain barrier. J Neuroimmunol 128: 58–68.
4. Fisher DT, Varanam TD, Muhitch JB, Evans SS (2010) Fine-tuning immune surveillance by fever-range thermal stress. Immunol Res 46: 177–189.
5. Turner GD, Morrison H, Jones M, Davis TM, Looverenwus S, et al. (1994) An immunohistochemical study of the pathology of fatal malaria. Evidence for widespread endothelial activation and a potential role for intercellular adhesion molecule-1 in cerebral sequestration. Am J Pathol 145: 1057–1069.
6. Staunton DE, Merluzzi VJ, Rothlein R, Barton K, Marlin SD, et al. (1989) A cell adhesion molecule, ICAM-1, is the major surface receptor for rhinoviruses. Cell 56: 849–853.
7. Chen Q, Appenheimer MM, Muhitch JB, Fisher DT, Clancy KA, et al. (2009) Thermal facilitation of lymphocyte trafficking involves temporal induction of intravascular ICAM-1. Microcirculation 16: 145–158.
8. Bielinski SJ, Pankow JS, Li N, Hsu FC, Adar SD, et al. (2008) ICAM1 and VCAM1 polymorphisms, coronary artery calcium, and circulating levels of soluble ICAM-1: the multi-ethnic study of atherosclerosis (MESA). Atherosclerosis 201: 339–344.
9. Fernandez-Reyes D, Craig AG, Kyes SA, Peshu N, Snow RW, et al. (1997) A high frequency African coding polymorphism in the N-terminal domain of the ICAM-1 predisposing to cerebral malaria in Kenya. Hum Mol Genet 6: 1357–1360.
10. Bellamy R, Kwiatkowski D, Hill AV (1998) Absence of an association between intercellular adhesion molecule 1, complement receptor 1 and interleukin 1 receptor antagonist gene polymorphisms and severe malaria in a West African population. Trans R Soc Trop Med Hyg 92: 312–316.
11. Kun NF, Klabunde JF, Del B, Luckner D, Alpers M, et al. (1999) Association of the ICAM-1 Klf1 mutation with protection against severe malaria in Lambaréne, Gabon. Am J Trop Med Hyg 61: 776–779.
12. Fry AE, Auburn S, Diakite M, Green A, Richardson A, et al. (2008) Variation in the ICAM1 gene is not associated with severe malaria phenotypes. Genes Immun 9: 462–469.
13. Register TC, Burdon KP, Lenchik L, Bowden DW, Hawkins GA, et al. (2004) Variability of serum soluble intercellular adhesion molecule-1 measurements attributable to a common polymorphism. Clin Chem 50: 2183–2187.
14. Gross MD, Bierlini SJ, Suarez-Lopez JR, Reiner AP, Bailey K, et al. (2012) Circulating soluble intercellular adhesion molecule 1 and subclinical atherosclerosis: the Coronary Artery Risk Development in Young Adults Study. Clin Chem 58: 411–425.
15. Waringe GM, Keane TM, Fegan G, Masyoki JN, Newton CR, et al. (2009) Plasmodium falciparum var gene expression is modified by host immunity. Proc Natl Acad Sci U S A 106: 21801–21806.
16. Bull FC, Paim A, Nhungo FM, Kanyanjui SM, Roberts DJ, et al. (2005) Plasmodium falciparum antigenic variation: relationships between in vivo selection, acquired antibody response, and disease severity. J Infect Dis 192: 1119–1126.
17. Cauwir J (1993) A Wilcoxon-type test for trend. Stat Med 4: 87–90.
18. Tchinda VH, Tadlen AD, Taklo EA, Tene G, Fogako J, et al. (2007) Severe malaria in Cameroonian children: correlation between plasma levels of three soluble inducible adhesion molecules and TNF-alpha. Acta Trop 102: 20–28.
19. Conroy AL, Phu H, Haukes M, Glover S, Mallesa M, et al. (2010) Endothelium-based biomarkers are associated with cerebral malaria in Malawian children: a retrospective case-control study. PLoS One 5: e15291.
20. Coerl-Gazdevich CM, Dzik WH, Erdman L, Sewanyana I, Dibangoye A, et al. (2010) Combined measurement of soluble and cellular ICAM-1 among children with Plasmodium falciparum malaria in Uganda. Malar J 9: 233.
21. Ashikpo S, Kusi KA, Ofori MF, Tetteh JK, Amoako-Sakyi D, et al. (2013) High Plasma Levels of Soluble Intercellular Adhesion Molecule (ICAM)-1 Are Associated with Cerebral Malaria. PLoS One 8: e44101.
22. Djoha Siawaya JF, Bapela NB, Ronacher K, Veenstra H, Kidd M, et al. (2008) Immune parameters as markers of tuberculosis extent of disease and early prediction of anti-tuberculosis chemotherapy response. J Infect Dis 56: 340–347.
23. Park GS, Ireland KF, Ojepka RO, John CC (2012) Evidence of Endothelial Activation in Asymptomatic Parasitemia and Effect of Blood Group on Levels of von Willebrand Factor in Malaria. J Pediatric Infect Dis Soc 1: 16–25.
24. Jenkins NE, Mwangi TW, Kortok M, Marsh K, Craig AG, et al. (2005) A polymorphism of intercellular adhesion molecule-1 is associated with a reduced incidence of nonmalarial febrile illness in Kenyan children. Clin Infect Dis 41: 1817–1819.
25. Mita-Mendoza NK, van de Hoef DL, Lopez-Mesa TM, Dombia S, Konate D, et al. (2013) A potential role for plasma uric acid in the endothelial pathology of Plasmodium falciparum malaria. PLoS One 8: e54481.
26. El-Deek SE, Makhlouf HA, Saleem TH, Mandour MA, Mohamed NA (2013) Sulfactant protein d, soluble intercellular adhesion molecule-1 and high-sensitivity C-reactive protein as biomarkers of chronic obstructive pulmonary disease. Med Princ Pract 22: 469–474.