NCF4 dependent intracellular reactive oxygen species regulate plasma cell formation

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ABSTRACT
Defective reactive oxygen species (ROS) production by genetically determined variants of the NADPH oxidase 2 (NOX2) complex component, NCF4, leads to enhanced production of autoantibodies to collagen type II (COL2) and severe collagen-induced arthritis (CIA) in mice. To further understand this process, we used mice harboring a mutation in the lipid endosomal membrane binding site (RS8A) of NCF4 subunit. This mutation did not affect the extracellular ROS responses but showed instead decreased intracellular responses following B cell stimulation. Immunization with COL2 led to severe arthritis with increased antibody levels in Ncf4 RS8A mutated animals without significant effects on antigen presentation, autoreactive T cell activation and germinal center formation. Instead, plasma cell formation was enhanced and had altered CXCR3/CXCR4 expression. This B cell intrinsic effect was further confirmed with chimeric B cell transfer experiments and in vitro LPS or CD40L with anti-IgM stimulation. We conclude that NCF4 regulates the terminal differentiation of B cells to plasma cells through intracellular ROS.

1. Introduction
Reactive oxygen species (ROS) and its downstream metabolites are known for their roles not only as toxic agents but also as physiological regulators of biological processes. The main source of ROS is the NOX2 complex [1]. The NOX2 complex is composed of a transmembrane heterodimer, NOX2 (gp91phox, CYBB) and p22phox (CYBA), a cytosolic heterotrimere, NCF1 (p47phox), NCF2 (p67phox) and NCF4 (p40phox) as well as a small GTPase Rac (1 or 2) [2]. Upon activation, phosphorylation of NCF1 induces conformational changes, allowing the complexing of NCF2 with the transmembrane heterodimer and Rac, which enables the transfer of electrons from NADPH to molecular oxygen to produce superoxide anion, and leads to hydrogen peroxide (H2O2) production as well as other kinds of ROS [3].

To anchor the complex, both NCF1 and NCF4 interact with membrane phospholipid through their PX domains, however, they have different preferences in phospholipid-binding specificities. NCF1 binds to phosphatidylinositol 3,4-bisphosphate and phosphatic acid [4,5], which are mainly found in plasma membrane, while NCF4 binds to phospholipid phosphatidylinositol 3-phosphate (PtdIns3P) [4], which is mainly found in endosome and phagosome membrane [6]. Therefore, NCF1 and NCF4 direct the NOX2 complex to different cellular compartments.

Deficiency in subunits of the NOX2 complex leads to chronic granulomatous disease (CGD), a rare primary immunodeficiency disease which is characterized by severe recurrent bacterial and fungal infections as well as tissue granuloma formation [7]. Compared to classical CGD, neutrophils from patients with mutations in NCF4 cannot kill bacteria efficiently but have an intact ability to kill fungi [8,9]. These NCF4-deficient patients suffer from hyperinflammation, autoimmunity and peripheral infections rather than life-threatening invasive infections. Interestingly, in vitro, NCF4 deficiency impairs NADPH oxidase activity of B cells more severely, compared to mononuclear phagocytes.
## Nomenclature

| Acronym | Description |
|---------|-------------|
| APC     | antigen presenting cell |
| ASC     | antibody secreting cell |
| BCR     | B cell receptor |
| CGD     | chronic granulomatous disease |
| CIA     | collagen-induced arthritis |
| COL2    | collagen type II |
| DPI     | days post immunization |
| GC      | germinal center |
| GSK     | GSK2795039 |
| H₂O₂    | hydrogen peroxide |
| iLN     | inguinal lymph node |
| LLPC    | long-lived plasma cell |
| MHC II  | major histocompatibility complex region class II |
| NOX2    | NADPH oxidase 2 |
| PB      | plasmablast |
| PIL     | pristane-induced lupus |
| Ptdlns3P| phospholipid phosphatidylinositol 3-phosphate |
| RA      | rheumatoid arthritis |
| ROS     | reactive oxygen species |
| SLE     | systemic lupus erythematosus |
| SNP     | single nucleotide polymorphism |
| Th      | follicular helper T |

[8]. Importantly, a single nucleotide polymorphism (SNP) in NCF1 has recently been identified [10,11], which has a major influence on a range of autoimmune diseases, including systemic lupus erythematosus (SLE) [10,12] and rheumatoid arthritis (RA) [12]. The association was not detected by genome wide association studies as the NCF1 gene has not been sequenced due to a complicated structure with duplications. Nevertheless, a copy number variation in NCF1 has been found associated with RA [11]. In addition, it was also found that SNPs linked to the NCF4 gene was associated with RA [13,14]. These findings confirm previous discoveries in animal models, where NCF1 was found to be major gene controlling both arthritis [15,16] and lupus [17]. The NCF1 effects on murine lupus confirmed previous findings that a deletion of NCF2 leads to severe lupus in autoimmune prone mouse strains [18].

The NCF4 protein interacts with endosomal membranes rather than the plasma membrane, due to its binding specificity for PtdIns3P, and it has been shown to be involved in the activation of NOX2 complex through Fc receptors and phagocytosis [19]. The interaction with PtdIns3P is mainly mediated by the arginine on position 58 (R58) and less efficient activation of the NOX2 complex on endosomal membranes was observed with this mutation [20].

To address if the profound effects on autoimmune were restricted to the NCF1 component, we recently showed that deletion of NCF4, as well as a mutation blocking its interaction with phospholipids, was strongly associated with collagen-induced arthritis (CIA) [21]. The deletion of NCF4 impairs the function of both NCF1 and NCF2, as these proteins are complexed. However, cells from mice with the R58A amino acid replacement mutation did not change the levels of other NOX2 complex components, showing a decreased intracellular ROS response but an increased extracellular response [21]. Interestingly, Ncf4<sup>R58A</sup> had a more restricted effect when tested in autoimmune conditions, with no effect in psoriatic arthritis while severely exaggerated disease was found in models of rheumatoid arthritis, such as CIA. In addition, it had a profound effect on antibody levels without disturbing the T cell response to collagen type II (COL2), suggesting there should be a direct regulation on B cell functions.

In this study, we first confirmed that the expression of cytosolic NOX2 complex subunits in B cells was not influenced by the Ncf4<sup>R58A</sup> mutation. This mutation preferentially influenced intracellular ROS response after PMA stimulation and had a more profound effect in B cells than in neutrophils. In the CIA model, the Ncf4<sup>R58A</sup> variant significantly enhanced disease severity and autoantibody production upon single immunization with COL2. During priming stage, the Ncf4<sup>R58A</sup> variant enhanced antibody secreting cell (ASC) formation and maturation with limited effects on antigen presentation of COL2, auto-reactive T cell activation and germinal center formation. Furthermore, COL2-specific plasma cells with Ncf4<sup>R58A</sup> showed altered CXCR3/CXCR4 expression and preferentially migrated to inflamed synovial tissues rather than bone marrow, which likely contributed to disease chronicity.

## 2. Results

### 2.1. Intracellular, but not extracellular ROS induction, is modified by Ncf4<sup>R58A</sup> in B cells

The deletion of NCF4 protein in Ncf4<sup>−/−</sup> mice has been reported to impair the expression of NCF1 and NCF2. Consequently, Ncf4<sup>−/−</sup> cells had decreased intra and extracellular ROS production. In contrast, neutrophils with the R58A mutation in the Ptdlns3P binding site of NCF4 had an unaffected expression of cytosolic NOX2 complex subunits as well as decreased intracellular ROS responses and increased extracellular ROS responses after stimulation [21]. To determine the influence of NCF4 on ROS production in B cells, we first measured the expression of cytosolic NOX2 complex subunits in B cells modified by Ncf4<sup>R58A</sup> (Fig. 1A-C, Fig. S1A).

Next, ROS production of splenic B cells from B10.Q.Ncf4<sup>R58A/S8A</sup> mice and littermate B10.Q.Ncf4<sup>R58A/R58</sup>, B10.Q.Ncf4<sup>R58/R58</sup> mice was measured. ROS produced by neutrophils were measured simultaneously as positive controls, as they are the strongest NOX2 complex-derived ROS producers. After stimulation with PMA, both neutrophils and B cells with Ncf4<sup>S8A/S8A</sup> showed decreased intracellular ROS responses (Fig. 1D-G), whilst extracellular ROS was not affected (Fig. 1H). ROS production of neutrophils from B10.Q.Ncf4<sup>S8A/R58</sup> and B10.Q.Ncf4<sup>R58/R58</sup> mice was 65.2% ± 3.5% and 88.8% ± 3.3% of B10.Q.Ncf4<sup>R58/R58</sup> neutrophils respectively (Fig. 1E), while the ROS production of B10.Q.Ncf4<sup>R58A/S8A</sup> and B10.Q.Ncf4<sup>R58A/R58</sup> B cells was 29.4% ± 5.2% and 77.0% ± 7.3% of B10.Q.Ncf4<sup>R58A/R58</sup> B cells (Fig. 1G), therefore showing a dose effect of Ncf4<sup>S8A</sup> mutation on intracellular ROS production in both B cells and neutrophils. Importantly, the normalized effect in B cells was even more pronounced than in neutrophils, suggesting that NCF4 may have a strong ROS-mediated intracellular influence on B cell function.

### 2.2. Ncf4<sup>S8A</sup> enhances autoantibody production and disease severity in CIA

Ncf4<sup>S8A</sup> has previously been shown to increase anti-COL2 antibody responses and disease severity in CIA [21]. To further explore the role of NCF4, we modified the CIA protocol by inducing the disease with a single priming immunization, as a booster dose of heterologous COL2 tends to enhance the immune response to the immunogen rather than allowing an autoimmune response from eroded cartilage in the joints (Fig. 2A and B). With a single priming immunization, Ncf4<sup>S8A</sup> dramatically enhanced the disease severity as well as the susceptibility to CIA. Even a single copy of the mutation had an effect, which correlated well with intracellular ROS production level, suggesting a dose-dependent effect of the intracellular ROS response on disease severity. Interestingly, if we applied the standard protocol of CIA that includes a booster immunization by COL2 in IFA on the 35th day post immunization (DPI),...
a diminished effect of the Ncf4<sup>R58A</sup> mutation was seen (Fig. S2A and B).

To study the formation of autoreactive ASCs, we measured anti-COL2 antibody titers in circulating blood during priming (12 DPI), disease onset (20 DPI), established (35 DPI) and chronic (90 DPI) stages. (Fig. 2C). Limited anti-COL2 antibodies were detected during priming stage and no differences were found between groups, while titers increased dramatically in Ncf4<sup>58A/58A</sup> mice on 20 DPI, a time point just before the disease onset. The isotypes of anti-COL2 antibodies on 20 DPI influenced by Ncf4<sup>58A</sup> were mainly IgG1 and IgG2b (Fig. 2D). Later in the disease development, the differences in anti-COL2 antibody titers diminished. Furthermore, the titer of anti-COL2 antibody on 20 DPI correlated with the later disease severity on 90 DPI (Fig. 2E), while the antibody titer on 90 DPI did not (Fig. 2F), suggesting that immune responses happened during priming stage is of greater importance for disease development.

To investigate whether pathogenic antibodies were produced by ASCs in situ, we measured ASCs in knee synovium on 90 DPI. CD19<sup>+</sup>B220<sup>+</sup>CD138<sup>+</sup> mature plasma cells but not CD19<sup>+</sup>B220<sup>+</sup>CD138<sup>+</sup> plasmablasts could be found in synovium of both B10Q.Ncf4<sup>S58A/S58A</sup> and B10Q.Ncf4<sup>R58/R58</sup> mice (Fig. 2G), with an increase of anti-COL2 ASCs from mice with Ncf4<sup>S58A</sup> (Fig. 2H), and most of them were IgG2b and IgG1 ASCs (Fig. 2I). The bone marrow is a survival niche for long-lived plasma cells that can give rise to circulating antibodies. Therefore, we also checked bone marrow from immunized (90 DPI) mice but we observed no effect by the Ncf4<sup>R58A</sup> mutation on COL2-specific ASCs (Fig. 2J and K). Finally, B10Q.Ncf4<sup>58A/58A</sup> mice had more active arthritis even at the very late stage as 90 days after immunization, which could be both a cause and result of the increased number of ASCs in the synovium.

2.3. T cell activation during priming stage is not affected by Ncf4<sup>R58A</sup>

The increased serum levels of anti-COL2 antibodies in B10Q.Ncf4<sup>S58A/</sup>S58A mice were IgG class-switched, suggesting that the B cell response after COL2 immunization was dependent on B-T cell interactions. To understand the mechanism of increased numbers of ASCs in Ncf4<sup>58A/58A</sup> mice, we first checked if antigen presentation and T cell activation during early priming stage (12 DPI) was affected. The number and major histocompatibility complex region class II (MHC II) expression on classical antigen presenting cells (APCs) (B cells, macrophages and dendritic cells) in spleen and inguinal lymph nodes (iLNs) were comparable between immunized B10Q.Ncf4<sup>S58A/R58</sup> and B10Q.Ncf4<sup>S58A/S58A</sup> mice. (Fig. 3A–D). Presentation of the MHCII molecule A<sub>q</sub> restricted major peptide of COL2 (both GalHyk264 or non-modified K264) and antigen processing of COL2 protein by naïve splenocytes, was also not affected in the B10Q.Ncf4<sup>S58A/58A</sup> mouse strain (Fig. 3E and F). The number of total CD4<sup>+</sup> T cells and the frequency of memory/effector/naive T cells were comparable between immunized B10Q.Ncf4<sup>S58A/R58</sup> and B10Q.Ncf4<sup>S58A/S58A</sup> mice. Samples without staining the target protein (indicated as fluorescence minus one, FMO) were used as negative controls. (D–G) Intra-cellular PMA-stimulated ROS production of spleen neutrophils (D) and B cells (F) from naive B10Q. Ncf4<sup>R58/R58</sup>, B10Q.Ncf4<sup>S58A/R58</sup> and B10Q.Ncf4<sup>S58A/S58A</sup> mice. Relative ROS production of spleen neutrophils (E) and B cells (G) was calculated by (MFI<sub>PMA</sub>-MFI<sub>control</sub>)/MFI<sub>control</sub>, then normalized to the mean of B10Q.Ncf4<sup>R58/R58</sup> group (wt). Results of relative ROS production were pooled from two independent experiments. (H) Extracellular PMA-stimulated ROS production of spleen leukocytes from naive B10Q. Ncf4<sup>S58A/R58</sup> and B10Q.Ncf4<sup>S58A/58A</sup> mice. FMO, fluorescence minus one; MFI, mean fluorescence intensity; DHR, dihydrorhodamine; wt, wild type.
comparable between immunized B10Q.Ncf4R58/R58 and B10Q.Ncf458A/58A mice (Fig. 3G and H). Moreover, we observed no differences in recall responses of Th1, Th2 and Th17 cells to the rat COL2 peptide Gal-HyK264 (Fig. 3I). Taken together, Ncf4R58A had no detectable influence on COL2 processing and presentation, or on antigen specific T cell activation.

2.4. Ncf458A enhances ASC formation with minor effects on germinal center formation

Next, B cell activation and differentiation during late priming stage (day 17 after COL2 immunization) was determined in the draining inguinal lymph nodes of immunized B10Q.Ncf4R58/R58 and B10Q.Ncf458A/58A mice. The total number of B cells, or expression of the activation marker CD86, was not affected by Ncf458A (Fig. 4A and B). However, the number of ASCs was clearly increased in B10Q.Ncf458A/58A mice on 90 DPI (Fig. 4F). Germinal center (GC) B cells showed an increased trend in terms of frequency in B10Q.Ncf458A/58A mice but no significant difference in the cell number (Fig. 4C) and comparable frequencies of dark zone and light zone GC B cells were observed (Fig. 4D). Accordingly, the number of follicular helper T (Tfh) cells was not affected either (Fig. 4E). ASCs were further subtyped: CD19 and B220 were used to distinguish plasmablast (PB)(B220+), newly formed plasma cell (CD19+B220+) and resting non-dividing plasma cell (or long-lived plasma cell [22], LLPC) (CD19+B220+) (Fig. 4H). Among all the ASCs, approximately 50% were LLPCs, 30% were newly formed PCs and 10–20% were PBs (Fig. 4H). Regarding the frequency of each subpopulations in total ASCs, B10Q.Ncf4R58/R58 mice had more
plasmablasts and B10Q.Ncf4R58A/S8A mice had more LLPCs (Fig. 4H). While in terms their frequency in live cells and their cell numbers, differences were not found in PBs but only in mature PCs (Fig. 4I and J), suggesting an increase of ASCs in Ncf4R58A/S8A mice mainly derived from B220+ mature PCs but not PBs. Antigen specific ASCs were also quantified and there were more anti-COL2 IgG ASCs in B10Q.Ncf4R58A/S8A mice (Fig. 4K). Interestingly, during early priming stage (12DPI), there were more anti-COL2 IgG and IgG2b ASCs in B10Q.Ncf4R58A/S8A mice (Fig. 4L) but no significant difference was found in general ASCs in iLN (Fig. 4G), suggesting that the role of NCF4 in preventing differentiation is stronger in autoreactive B cells than in other non-autoreactive B cells. ASCs and anti-COL2 ASCs could also be detected in bone marrow and spleen of COL2-immunized mice (17 DPI), but no differences were observed between B10Q.Ncf4R58A/S8A and B10Q.Ncf4S8A/S8A group (Fig. S3A-D). Similar increases were also present in the pristane-induced lupus (PIL) model. Mice received a single i.p. injection of pristane, resulting in more anti-dsDNA and anti-nucleosome IgG plasma cells in bone marrow from immunized B10Q.Ncf4R58A/S8A mice compared to B10Q.Ncf4R58R/S8R mice (Fig. 4O), but with relatively unaffected number of GC B cells and Tfh cells (Fig. 4M, N) on the 10th day post injection. This suggests that the observed effect by Ncf4R58A on ASC formation was not restricted to COL2. Taken together, the Ncf4R58A mutation could enhance PC formation and autoantibody secretion, but with no or only minor effects on GC formation.

2.5. ASC formation regulated by Ncf4R58A is a B cell intrinsic effect

To investigate whether the enhanced ASC formation in COL2-immunized B10Q.Ncf4R58A/S8A mice is a B cell intrinsic or extrinsic effect, we performed B cell transfer experiments. B cells sorted from male B10Q.Cd45R2.Ncf4S8A/S8A (all leukocytes express CD45.2) and B6NQ.Cd45R1 mice were equally mixed and transferred to B10Q.Cd45R2 μMt.Ncf4S8A/S8A mice, which lack mature B cells. Recipient mice were
Fig. 4. Ncf4<sup>R58A</sup> enhanced ASC formation with minor effects on germinal center formation
(A) The frequency, number and (B) CD86 expression of B cells in iLN from immunized B10.Q.Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice on 17 DPI. Results were pooled from two independent experiments. (C) The frequency and number of germinal center B cells (GL7<sup>+</sup>CD95<sup>+</sup> cells, gate on CD3<sup>+</sup>CD4<sup>+</sup> cell population) in iLN from naïve and immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice on 17 DPI. Representative gate was shown. (D) The frequency of dark zone (CXCR4<sup>+</sup>CD86<sup>+</sup>) and light zone (CXCR4<sup>+</sup>CD86<sup>+</sup>) GC B cells in iLN from immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice on 17 DPI. Representative gate was shown. (E) The frequency and number of follicular helper T (Tfh) cells (CXCR5<sup>+</sup>PD1<sup>+</sup> cells, gate on CD3<sup>+</sup>CD4<sup>+</sup> cell population) in iLN from immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice (17 DPI). Representative gate was shown. (F) The frequency and number of ASCs (CD138<sup>+</sup>Sca-1<sup>+</sup>IgD<sup>+</sup>) in iLN from naïve and immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice on 17 DPI. Representative gate was shown. (G) The frequency and number of ASC subpopulations: plasmablasts (PBs) (CD19<sup>+</sup>B220<sup>+</sup>), newly formed plasma cells (NEW PCs) (CD19<sup>+</sup>B220<sup>-</sup>) and long-lived plasma cells (LLPCs) (CD19<sup>-</sup>B220<sup>-</sup>) in total ASCs (H) and live cells (I) of iLN from immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice on 17 DPI. Representative gate was shown. (J) The number of PBs, NEW PCs and LLPCs in iLN from immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice (17 DPI). (K) Total anti-COL2 IgG ASCs and anti-COL2 IgG2b ASCs in iLN from immunized B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice on 12 DPI. Representative anti-COL2 IgG2b ELISPOT results were shown. (M) Mice were induced lupus using a single i.p. injection of pristane (pristane-induced lupus, PIL). The frequency and number of germinal center B cells (GL7<sup>+</sup>CD95<sup>+</sup> cells, gate on CD3<sup>+</sup>CD4<sup>+</sup> cell population) in spleen from B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice were checked on 10 days post injection (dpi). (N) The frequency and number of Tfh cells (CXCR5<sup>+</sup>PD1<sup>+</sup> cells, gate on CD3<sup>+</sup>CD4<sup>+</sup> cell population) in spleen from B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice in PIL model (10 dpi). (O) Total IgG ASCs, anti-dsDNA ASCs and anti-nucleosome ASCs in bone marrow and spleen from B10.Q. Ncf4<sup>R58R/R58R</sup> and B10.Q. Ncf4<sup>S6A/S6A</sup> mice in PIL model (10 dpi). GC B, germinal center B cells; Tfh, follicular helper T; PB, plasmablast; NEW PC, newly formed plasma cell; LLPC, long-lived plasma cell; PIL, pristane induced lupus; dpi, days post injection.
immunized with bovine COL2, 2 days after transfer (day 0) (Fig. 5A). Purity of B cells after sorting and transfer efficacy were checked on day –2, day 0 and 16 DPI (Fig. S4A-D). Spleen, lymph nodes and bone marrow were harvested 16 days after immunization. B cells were present 2, day 0 and 16 DPI (Fig. S4A-D). Spleen, lymph nodes and bone marrow, and no differences in ILN, mesenteric lymph nodes and blood (Fig. S4F). The majority of transferred B cells stayed in the spleen (Fig. S4E), possibly because they were transferred intravenously. We next measured GC and ASC formation in these mice. Most ASCs were derived from B10Q.Cd45.7, Ncf4<sup>58A/58A</sup> B cells while more GCs were formed from B6N.Cd45.1 B cells (Fig. 5B and C), suggesting that NCF4 had a B cell intrinsic effect in both activation and differentiation. The ASC population was further divided into PB (9%), newly formed PC (27%) and LLPC (60%) (Fig. 5D), and the influence of Ncf4<sup>58A/58A</sup> on ASC formation was mainly found in newly formed PC (CD19<sup>+</sup>B220<sup>+</sup> ASC) (Fig. 5E and F).

It has been reported that, CD45.1/Cd45.2 congenic bone marrow transplantation has a sex-related reconstitution bias [23, 24] as CD45.1<sup>+</sup> B cells had a reduced reconstitution potential in female recipients. We also observed a related phenomenon in our experiments, despite the fact that we used mature B cells for transfer. When we checked B cells in the peripheral blood, spleen, lymph nodes and bone marrow 2, day 0 and 16 DPI (Fig. S4E-G) on 16 DPI. There were slightly more CD45.1<sup>+</sup> B cells in spleen, axillary lymph nodes and bone marrow compared with mice transferred with B10Q.Cd45.7, Ncf4<sup>58A/58A</sup> mice, and CD45.2<sup>+</sup> B cells than B10Q.Cd45.7, Ncf4<sup>58A/58A</sup> mice while more GCs were formed from B6N.Cd45.1 B cells (Fig. 5E and F), confirming the difference was dependent on Ncf4<sup>58A/58A</sup>.

Next, we checked if these PCs, formed after transfer, functioned normally. This time B10Q.Ncf4<sup>58A/58A</sup> B cells and B10Q.Ncf4<sup>58A/58A</sup> B cells were transferred separately into B10Q.Cd45.7, Ncf4<sup>58A/58A</sup> mice and immunized 2 days later (Fig. 5H). There were more anti-COL2 IgG antibodies on 20 DPI and 42 DPI in mice transferred with B10Q.Ncf4<sup>58A/58A</sup> B cells, compared with mice transferred with B10Q.Ncf4<sup>58A/58A</sup> B cells (Fig. 5I).

Taken together, these data showed that B cells expressing Ncf4<sup>58A/58A</sup> were more competitive in forming ASCs.

Fig. 5. B cells with Ncf4<sup>58A</sup> were more competitive in forming ASCs. (A) Scheme of B6N.Cd45.1<sup>+</sup> B10Q.Cd45.2<sup>+</sup>Ncf4<sup>58A/58A</sup> B cell transfer experiment. (B) Representative plots and (C) percentage of CD45.1<sup>+</sup> and CD45.2<sup>+</sup> cells in B cells (CD19<sup>+</sup>B220<sup>+</sup>), GC B cells (CD95<sup>+</sup>GL7<sup>+</sup>, gate on IgD<sup>-</sup> B cells) and ASCs (CD138<sup>+</sup>Sca-1<sup>+</sup>, gate on IgD<sup>-</sup>) in the spleen of immunized recipient μMt.Ncf4<sup>58A/58A</sup> mice on 16 DPI. Immunized μMt.Ncf4<sup>58A/58A</sup> mice without transferring B cells were used as negative controls for each gating. (D) Representative plots and percentage of ASC subpopulations: PB (CD19<sup>+</sup>B220<sup>+</sup>), NEW PC (CD19<sup>+</sup>B220<sup>+</sup>) and LLPC (CD19<sup>+</sup>B220<sup>+</sup>) in ASCs in the spleen of immunized recipient μMt.Ncf4<sup>58A/58A</sup> mice on 16 DPI. (E) Frequency of CD45.1<sup>+</sup> and CD45.2<sup>+</sup> cells in PB, PB, NEW PC and LLPC in the spleen of immunized recipient μMt.Ncf4<sup>58A/58A</sup> mice on 16 DPI. (F) Frequency of CD45.1<sup>+</sup> PB, PB, NEW PC, LLPC and CD45.2<sup>+</sup> PB, PB, NEW PC, LLPC in live cells in the spleen of immunized recipient μMt.Ncf4<sup>58A/58A</sup> mice on 16 DPI. (G) Percentage of CD45.1<sup>+</sup> and CD45.2<sup>+</sup> cells in ASCs (CD138<sup>+</sup>Sca-1<sup>+</sup>, gate on IgD<sup>-</sup>) cells) in the spleen of immunized μMt.Ncf4<sup>58A/58A</sup> mice transferred with B10Q.Cd45.7, Ncf4<sup>58A/58A</sup> or B10Q.Cd45.2<sup>+</sup>Ncf4<sup>58A/58A</sup> mice on 16 DPI. (H) Scheme of B10Q.Ncf4<sup>58A/58A</sup> and B10Q.Ncf4<sup>58A/58A</sup> B cell transfer experiment. (I) Levels of anti-COL2 antibodies in the serum from immunized recipient μMt.Ncf4<sup>58A/58A</sup> mice transferred with B10Q.Ncf4<sup>58A/58A</sup> B cells or B10Q.Ncf4<sup>58A/58A</sup> B cells on day 0, 20 DPI and 42 DPI.

preference for differentiation to functional plasma cells secreting autoreactive anti-COL2 antibodies.

### 2.6. Differentiation of Ncf48A-expressing B cells to plasmablasts is promoted by their lower intracellular ROS level

We established an *in vitro* system to identify the conditions whereby Ncf48A could enhance the PC differentiation. We cultured B cells for 3 days with various known stimuli (LPS, CD40L i.e., CD40L expressing fibroblasts, IL21+CD40L, anti-μ+IL21+CD40L, IL4+IL21+CD40L or IL4+IL21+anti-μ+CD40L) with no effect on cell viability between Ncf48A/- and Ncf48A/8A groups (Fig. S5A). After 3 days LPS stimulation, B cells from B10Q.Ncf48A/- spleen clearly formed more plasmablasts both in terms of frequency and cell number (Fig. 6 A). There were also more IgM in the supernatant of cultures with B10Q.Ncf48A/- B cells (Fig. 6B). In the CD40L stimulating group, the results were however not consistent. When stimulating with CD40L, IL21, with or without (w/wo) IL4, no difference was found between B10Q.Ncf48A/- and B10Q.Ncf48A/8A B cells. However, if we added anti-μ stimulation in this system, we found more plasmablasts formed from B cells with Ncf48A/8A (Fig. 6C). Because anti-μ antibody neutralized Ig secretion, we could not observe antibody differences in the supernatant (Fig. 6D). These results suggest that Ncf48A promotes ASC formation if stimulated through the TLR4 or the B cell receptor (BCR) pathway.

Next, we determined if stimulating TLR4 or BCR could activate the NOX2 complex and if it was NCF4-dependent. Intracellular ROS production of stimulated B cells was measured by dihydrorhodamine (DHR) assay. We found that LPS activation of B cells with CD40L, anti-μ, IL4 or CD40L+IL4+anti-μ led to a lower ROS response than Ncf48A/8A B cells (Fig. 6F). Although CD40 ligation was reported to be able to induce ROS burst through the NOX2 complex [25], ROS production stimulated by CD40L+IL21 w/wo IL4 was not affected by Ncf48A. However, stimulation through the B cell receptor induced intracellular ROS production at a lower level in Ncf48A/8A B cells (Fig. 6E), similar to the LPS stimulation (Fig. 6F).

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Fig. 6. Differentiation of Ncf48A-expressing B cells to plasmablasts *in vitro* was promoted by their lower intracellular ROS level

(A) The number and frequency of plasmablasts within B10Q.Ncf48A/8A and B10Q.Ncf48A/- B cells after stimulated with LPS for 84 h. Representative gate was shown. (B) Secreted IgM in the culture supernatant of B10Q.Ncf48A/- and B10Q.Ncf48A/8A B cells after stimulated with LPS for 84 h. (C) The number and frequency of plasmablasts within B10Q. Ncf48A/8A and B10Q.Ncf48A/- B cells after stimulated with irradiated CD40L expressing fibroblasts (CD40L), CD40L+IL21; CD40L+IL21+anti-μ; CD40L+IL21+IL4 or CD40L+IL21+IL4+anti-μ for 84 h. Representative gates were shown. (D) Secreted IgM in the culture supernatant of B10Q.Ncf48A/- and B10Q.Ncf48A/8A B cells after stimulated with CD40L, CD40L–IL21; CD40L–IL21+anti-μ; CD40L+IL21+IL4 or CD40L+IL21+IL4+anti-μ for 84 h. (E) Intracellular ROS production of B10Q. Ncf48A/8A and B10Q.Ncf48A/- B cells after stimulated by CD40L+IL21; CD40L+IL21+anti-μ; CD40L+IL21+IL4 or CD40L+IL21+IL4+anti-μ for 1 day. (F) Intracellular ROS production of B10Q. Ncf48A/8A and B10Q.Ncf48A/- B cells stimulated by LPS and LPS + GSK2795039 (GSK) for 1 day. (G) The number and frequency of plasmablasts within B10Q.Ncf48A/8A and B10Q.Ncf48A/- B cells stimulated with LPS in the presence of GSK or H2O2 (during 40 h–84 h of incubation). GSK, GSK2795039.
To confirm that this effect on differentiation was ROS dependent, we added the NOX2 complex inhibitor GSK2795039 (GSK) or H2O2 to LPS-stimulated B cells at the later differentiation stage (40 h–84 h of incubation) [26]. As shown in Fig. 6F, GSK could inhibit ROS production from NOX2 complex effectively. After 3 days incubation, there were more plasmablasts formed in both Ncf4<sup>R58/R58</sup> and Ncf4<sup>58A/58A</sup> groups inhibited with GSK (Fig. 6G). As Ncf4<sup>58A</sup> only partly decreased ROS production, a further decrease of intracellular ROS by GSK enhanced plasmablasts formation further. After adding H2O2, the number of plasmablasts decreased in both groups and no differences were observed between Ncf4<sup>R58/R58</sup> and Ncf4<sup>58A/58A</sup> groups anymore (Fig. 6G). These results confirmed that the regulation by Ncf4<sup>58A</sup> on ASC formation operated through intracellular ROS production.

2.7. Altered expression of CXCR3 and CXCR4 on ASCs from B cells with Ncf4<sup>58A</sup>

The increased number of ASCs formed in draining lymph node follicles of Ncf4<sup>58A/58A</sup> mice migrated preferentially to the inflamed joint synovial tissues and we next attempted to explain this migration pattern.

We observed a decrease in the number of ASCs (Fig. S6A) and almost no PBs (Fig. S6C) in iLN on 20 DPI compared to that on 17 DPI (Fig. S6B), suggesting that the priming B cell responses were mostly completed and ASCs had migrated out of iLNs before 20 DPI. Next, we performed chimeric B cell transfer experiments (performed in Fig. 5A), CXCR4 was expressed higher level of CXCR4 while this effect was gradually diminished as B cell differentiated to PBs and PCs. Notably, the long-lived PCs (CD19<sup>B220</sup>) with Ncf4<sup>58A/58A</sup> showed lower expression of CXCR4 compared to plasmablasts derived from B10Q. Ncf4<sup>R58/R58</sup> and B10Q.Ncf4<sup>58A/58A</sup> B cells, as CXCR4 homes ASCs to bone marrow, while CXCR3 drives ASCs to inflamed tissues [27]. In the chimeric B cell transfer experiments (performed in Fig. 5A), CXCR4 was upregulated during B cell differentiation and ASC maturation (Fig. 7A). B cells with Ncf4<sup>58A/58A</sup> expressed higher level of CXCR4 while this effect was gradually diminished as B cell differentiated to PBs and PCs. By studying the effect of a mutation (R58A) on Ncf4 which results in a replacement of an amino acid interacting with endosomal lipid membranes, we could investigate the role of intracellular ROS on B cell activation. We found that the decreased intracellular ROS due to the Ncf4<sup>58A</sup> mutation did not affect interactions with T cells but could promote ASC formation and maturation to plasma cells. These cells produced autoreactive COL2-specific antibodies and preferentially migrated to inflamed synovial tissues rather than bone marrow, where likely contributed to sustained disease and chronicity.

Ncf4 (alternatively named p40phox) is a cytosolic protein which together with NCF1 and NCF2 adheres to the membrane NOX2 complex component to induce a ROS response. A role of NCF4 is to preferentially direct the cytosolic NCF1-NCF2-NCF4 proteins to endosomal membranes to activate the NOX2 complex intracellularly. An arginine

3. Discussion

To clarify the importance of genetic polymorphisms associated with NOX2 complex in B cell dependent autoimmune diseases requires a greater understanding of how NOX2 complex-derived ROS regulate B cells. By studying the effect of a mutation (R58A) on Ncf4 which results in a replacement of an amino acid interacting with endosomal lipid membranes, we could investigate the role of intracellular ROS on B cell activation. We found that the decreased intracellular ROS due to the Ncf4<sup>58A</sup> mutation did not affect interactions with T cells but could promote ASC formation and maturation to plasma cells. These cells produced autoreactive COL2-specific antibodies and preferentially migrated to inflamed synovial tissues rather than bone marrow, where likely contributed to sustained disease and chronicity.

Ncf4 (alternatively named p40phox) is a cytosolic protein which together with NCFI and NCF2 adheres to the membrane NOX2 complex component to induce a ROS response. A role of NCF4 is to preferentially direct the cytosolic NCF1-NCF2-NCF4 proteins to endosomal membranes to activate the NOX2 complex intracellularly. An arginine
replacement to an alanine at position 58 affects NCF4 adhesion to phospholipid phosphatidylinositol 3-phosphate (PtdIns3P) [28] resulting in reduction of intracellular ROS responses, whereas knockout of NCF4 protein reduces NCF1 and NCF2 components expression leading to blockage of both intra and extracellular ROS responses. The m1j mutation in Ncf1, which results in a non-functional truncated protein and blocking of the lipid membrane adhesion, also reduces both intra and extracellular ROS responses, and has a profound effect on tolerance and autoimmune diseases [29]. Thus, the Ncf4<sup>R58A</sup> mutation provides a unique opportunity to understand the effects of intracellular ROS responses by the NOX2 complex.

Previously, we have shown that ROS produced by antigen presenting cells could oxidize thiols on and inside T cells [30], thereby protect against autoimmune arthritis models. Such a mechanism is dependent on NCF1 while the observation here suggests that NCF4, affecting intracellular ROS only, cannot mediate such protective effects on T cells in CIA. However, human NCF4-deficient (Ncf4<sup>R58G</sup>) B cells displayed a reduced capacity of presenting several exogenous antigens while presentation of membrane autoantigens was efficient [31]. Taken together, the influence of NCF4 on antigen presentation differs by antigen specificity and NCF4 may skew the epitope selection.

Where exactly the activation of the NOX2 complex happens during B cell differentiation is however not clear. As the activation requires interaction with antigens, there are several phases where this could occur: extracellular interaction with antigens, the interaction with T cells at T-B border, and the repetitive activation during the germinal center reaction. All these interactions can lead to the differentiation towards plasmablasts. Plasmablasts from an extracellular interactions typically peak 4–6 days after immunization while plasmablasts produced from the GC response peaks 2 weeks after immunization [32]. In our study, we observed an increase in PC formation at the priming stages accompanied with sustained higher isotype-switched anti-COL2 antibody titers in mice with Ncf4<sup>R58A</sup>. As the half-life of most plasmablasts is only a few days [33], the observation that cells secreting anti-COL2 IgG in the synovial tissues 90 days after primary immunization most likely reflects that they ended up as long-lived plasma cells secreting pathogenic antibodies in the targeted cartilaginous joints. Studies have suggested that long-lived plasma cells are mostly derived from an authentic germinal center reaction [32]. Thus, most likely, the Ncf4<sup>R58A</sup> mutation also promotes plasma cell formation from the germinal center reaction.

The migration of plasma cells to the joints is compatible with the prolonged chronic arthritis seen in the Ncf4<sup>R58A</sup> mutated mice. It is well recognized that chronic inflammation could lead to modulation of chemokine receptor expression on B cells and plasma cells [34,35]. CXCR3 expression of B cells and plasma cells is up-regulated in rheumatoid arthritis and systemic lupus erythematosus [36]. However, our study showed that CXCR3 expression on B cells and ASCs could be regulated intrinsically before the inflammation occurred. NCF4-dependent ROS induced by BCR ligation is needed for controlling the expression of CXCR3, indicating another aspect of redox regulation on B cell differentiation.

The COL2-B cells are autoreactive. They secrete pathogenic antibodies binding cartilage in vivo. Nevertheless, COL2-B cells are positively selected in the bone marrow, which was confirmed here by the presence of these cells in healthy mice [37]. It means that COL2 autoreactive B cells are already selected and present at the time of the COL2 immunization. Indirect evidence argues for that such B cells are regulatory [37], but if they differentiate to plasma cells, they can secrete pathogenic antibodies. It is possible that ROS induced during cell activation in the follicle, hinder the cells to differentiate to plasma cells and therefore protecting from disease enhancement. This could be generalized to all autoreactive B cells with a potential to produce autoantibodies, both natural polyreactive antibodies, as in lupus, and more specific cartilage specific antibodies, as in arthritis. To drive B cells to plasma cells we immunized the mice with rat or bovine COL2, which activates COL2-reactive T cells, thereafter activating COL2-reactive B cells. Decreased intracellular ROS in Ncf4<sup>R58A</sup> COL2-reactive B cells lowered the threshold for differentiating to plasma cells which could secrete pathogenic antibodies, leading to arthritis.

The oxygen metabolites that mediates signaling from the ROS induction is hydrogen peroxide, due to its sustained stability, which can oxidize selected sets of thiols on proteins and signaling pathways of the B cells [38]. There are however 20–40,000 such thiols in a B cell and it is likely that numerous interacting pathways can be affected [39], reaching a high level of complexity. BCR signaling plays a central role in controlling B cell fate decisions. Depending on the context, BCR ligation could lead to cell survival, apoptosis, proliferation or differentiation [40]. ROS has been shown to augment BCR signaling by inhibiting the activity of protein tyrosine phosphatase which negatively regulates BCR signaling [41]. NOX2 complex is responsible for generating the rapid initial production of ROS [42] while BCR proximal signaling and downstream signaling pathways are not affected by NOX2 [43,44]. Bertolotti et al. showed that gp91<sup>phox</sup>−/− B cells had impaired IgM secretion upon LPS stimulation [38] possibly because an early oxidative step was necessary to start the differentiation program [26]. In late-differentiating B cells, antioxidant induced an increase in antibody production [42], suggesting NOX2-derived ROS play a dule role in different stages of B cell differentiation. However, exactly how NOX2-derived hydrogen peroxide targets different pathways can only be solved on a higher level of complexity involving interactions of thousands of redox regulated targets in different interacting pathways.

4. Conclusions

This study shows that B cells are regulated by intrinsically produced ROS, derived from the NOX2 complex and modified by NCF4. Mutations affecting an arginine at position 58 on NCF4 regulates intrinsic oxidative burst in B cells, driving plasma cells formation and influencing their migration patterns, which is a risk factor for the development of autoimmune manifestations.

5. Materials and methods

5.1. Mice

Founders of Ncf4<sup>R58A/R58A</sup> mice [28] were generously provided by Phillip T. Hawkins (Babraham Institute, Cambridge, UK) and crossed with C57BL/10.Q/rhd mice (abbreviated B10.Q.Ncf4<sup>R58A/R58A</sup>) for more than 10 generations to obtain an arthritis-permissive major histocompatibility complex type II (MHC II)<sup>A</sup> haplotype. B10.Q.Ncf4<sup>R58A/R58A</sup> mice were obtained from heterozygous breeding and cohoused with heterozygous and wild-type littermate controls (indicated as B10.Q.Ncf4<sup>R58A/R58A</sup> and B10.Q.Ncf4<sup>R58A/R58A</sup>). Cd45<sup>−/−</sup> mice (B6.SJL-Pgpc<sup>−/−</sup> Pepc<sup>−/−</sup>Boy1) and µMt<sup>−/−</sup> mice (B6.129S2-Ighm<sup>1Gμ1</sup>/J) were purchased from Jackson Laboratory. They were crossed with C57BL/6N.Q mice (abbreviated B6N.Cd45<sup>−/−</sup>) and B10.Q mice (abbreviated B1010.Q.µMt<sup>−/−</sup>) respectively for more than 10 generations to obtain the H2-C<sup>A</sup> haplotype. Then, B1010.Q.µMt<sup>−/−</sup> mice were crossed with B10.Q.Ncf4<sup>R58A/R58A</sup> mice to obtain B cell deficient mice with Ncf4<sup>R58A/R58A</sup> background (abbreviated µMt<sup>−/−</sup> Ncf4<sup>R58A/R58A</sup>).

Genotype of B10.Q.Ncf4<sup>R58A/R58A</sup> mice was determined by qPCR with a primer set of forward 5‘ CAAAGAGGAGGTCAATGATCCTCA 3’, reverse 5‘ CAAAACGGTCCTCGAGCCT 3’ and two specific dye-taged Taqman probe TACCGGCTATC [FAM] and TACCGGGCTATC [VIC]. Genotype of B10.Q.µMt<sup>−/−</sup> mice was determined by PCR with a primer set of 5‘ CCG TCT AGG TGG TAC TAT TAG G 3’, 5‘ GAA GAC GAT GAA GTG GGG 3’, and 5‘ TGG TGG CCA GTC ATC GGA 3’, which was re-confirmed that no B cells could be found in peripheral blood by staining anti-CD19, anti-B220 and checking by flow cytometry. Genotype of Cd45<sup>−/−</sup> mice was determined by CD45.1/CD45.2 expression on leukocytes in peripheral blood. White blood cells were stained with anti-CD45.1 and anti-CD45.2 and checked by flow cytometry.
All mice were kept under controlled temperature and humidity with a 12-h light-dark regimen environment in polystyrene cages (IVF) under specific pathogen free condition in the animal section of medical inflammation research at the Karolinska Institute, Stockholm. Experiments followed the ARRIVE guidelines [45].

5.2. Antibodies and flow cytometry

Antibodies were purchased from BD Bioscience, Biolegend, and EBioscience, including anti-CD19(1D3), anti-CD45R(B220)(RA3-6B2), anti-CD21/CD35(7E9), anti-CD3 (RM2-5), anti-CD93(AA4.1), anti-CD23(2G8), anti-CD21(D35.7E9), anti-CD3a(145-2C11), anti-CD4(M4-5), anti-CD62L (MEL-14), anti-CD44(UTM7), anti-CD8 (53-6.7), anti-CD11b(M1/70), anti-CD11c(N418), anti-Ly6C(AL-21), anti-Ly6G(1A8), anti-H-2, I-A/I-E(2G9), anti-F4/80(RM8), anti-CD138(28-1-2), anti-Ly6a/E(sca-1)(D7), anti-Ly6C(GL7), anti-CD95(15A7), anti-CD45.1(A20), anti-CD45.2 (104), anti-CD184(CXCR4)(1276F12), anti-CD86(GL1), anti-CD183 (CXCR3)(CXCR3-175), anti-CXCR5(CD185) (SPRCLS5), anti-CD279 (PD-1) (RMPI-30) with the fluorescence FITC, PE, PerCP-Cy5.5, PE-Cy7, APC, Alexa Fluor 700, APC-Cy7, Pacific Blue, BV605 or BV650. Dead cells were excluded by fixable near-IR dead cell stain kit (Molecular Probes, #L34976B). Briefly, for staining markers on cell surface, single cell suspension was obtained from spleen, lymph nodes and bone marrow. Spleen and bone marrow cells were first incubated with ammonium chloride-potassium (ACK) buffer (homemade) for 5 min at RT to lyse red blood cells. Then, cells were washed, incubated with Fcγ receptor blocker (24G2) for 10 min at RT followed by 20 min surface markers staining in dark on ice. For intracellular staining of NOX2 complex cytosolic subunits, cells were first stained with surface markers, fixed and permeabilized by Fixation and Permeabilization buffer (BD Bioscience, #554722). Then, they were stained with anti-p47phox (NCF1, D-10), anti-p67phox (NCF2, D-6) and anti-p40phox (NCF4, D-8) (all from Santa Cruz). Samples without staining the targeted protein, which were indicated as fluorescence minus one (MOM), were used as the negative controls. After that, cells were washed, acquired with Attune NxT flow cytometer (ThermoFisher) and analyzed by Flowjo.10.6.

5.3. ROS detection

Intracellular ROS was detected by flow cytometry. Briefly, cells were first stained with cell surface markers, then incubated with 3 μM dihydorhodamine (DHR) 123 (Invitrogen, #D23806) for 10 min at 37 °C and stimulated with 200 ng/ml PMA (Sigma-Aldrich, #P1585) for 20 min at 37 °C. Dimethyl sulfoxide (DMSO) (Molecular Probes, #L34976B) was used for control group. Geometric mean fluorescence intensity (MFI) of DHR was detected by flow cytometry.

Extracellular ROS production was detected by chemiluminescence assay. Briefly, cells were stimulated with 100 ng/ml PMA for 60 min in HBSS (with Ca²⁺ and Mg²⁺ -) (ThermoFisher, #14025050) containing 150 μM isoluminol (Sigma-Aldrich, #82624) and 18.75U/ml HRP II (Sigma-Aldrich, #P8250). Cells without PMA stimulation were used as negative controls. Data output was measured in relative light units (RLU).

5.4. Induction and evaluation of collagen-induced arthritis (CIA)

Rat type II collagen (COL2) was obtained from pepsin-digested SWARM chondrosarcoma, and subsequently processed as previously described [46]. Mice were immunized with rat COL2 (1 mg/ml, 100 mg per mouse) emulsified 1:1 in 50 μl complete Freund’s adjuvant (CFA, Nordic Biolabs #263810) intradermally at the base of tail. When indicated in the figure, they were boosted on the 35th day after primary immunization with rat COL2 emulsified 1:1 in incomplete Freund’s adjuvant (IFA, Nordic Biolabs #263910) in a total volume of 50 μl. All experiments were performed blindly, with age and sex-matched littersmates and with groups randomly distributed in cages.

Arthritis severity was scored using a macroscopic scoring system [47] blindly at the indicated time points after primary immunization. Briefly, 1 point was given for one red and swelling finger/toe, 1–5 points were given for each inflamed ankle/wrist or paw/hand according to the severity. A total of 15 points could be given to one limb and all mice were scored arrange from 0 to 60 points. Mice were bled from the sub-mandibular vein at indicated time points in order to detect anti-COL2 antibodies by ELISA. Briefly, serum was diluted and added to COL2-coated (10 μg/ml) ELISA plates (Biolegend, #423501). Bound IgGs were detected with HRP-conjugated goat anti-mouse Ig kappa (Southern Biotech, #1170-05) or anti indicated isotypes of mouse IgG (anti-IgG1 [1070-05], anti-IgG2b [1091-05], anti-IgG2c [1078-05], anti-IgM [1021-05], all from Southern Biotech) and ABTS (Roche, #11112422001). The absorbance was read at 405nm (Synergy-2; BioTek Instruments). For total anti-COL2 antibodies, recombinant anti-COL2 antibody (0.1 mg/ml, homemade) was used as standard. All animal experiments were performed under valid ethical permits approved by animal welfare authorities (Stockholm region, Sweden).

5.5. Antigen-specific antibody secreting cells detection

Single cell suspension was obtained from spleen, lymph nodes, bone marrow, synovium and resuspended in complete RPMI (ThermoFisher, #61870044) media containing 10% FCS (ThermoFisher, #26140079) and penicillin/streptomycin (Sigma, #P4333). Then, cells were added into COL2-coated (10 μg/ml) dsDNA-coated (20 μg/ml) Sigma-Aldrich, #D3664), nucleosome-coated (10 μg/ml, homemade) or anti-IgG-coated (1 μg/ml) ELISPOT plates (Merck Millipore, #MISP54W10). For dsDNA coating, Poly-L-Lysine (20 μg/ml, #P2658) was pre-coated one day before and dsDNA was coated in sterile TE buffer. After 2 h incubation at 37 °C, plates were washed and detected by biotinylated goat anti-mouse IgG (Southern Biotech, #1030-08), IgG1 (Southern Biotech, #1070-08) or IgG2b (Southern Biotech, #1090-08), followed by ExtrAvidin® conjugated alkaline phosphatase (Sigma-Aldrich, #E2636) and BCIP/NBT (Sigma-Aldrich, #B5655). Spots were scanned with ImmunoScan and analyzed with ImmunoSpot software (Cellular Technology Ltd.).

5.6. T cell recall assay

Peptides spanning the sequence 259–273 of COL2 with a non-modified lysine at position 264 (K264) or with a β-galactopyranosyl residue on ε-hydroxylsine at position 264 (GalHyK264) were synthesized as previously described [48]. Cells from spleen and inguinal lymph nodes were subjected to anti-IFNγ (AN18) (Mabtech, #3321-3-1000), anti-IL17A (TC11-18H10.1) (BD, #555068), anti-IL-4 (11B1) (BD, #554394) (5 μg/ml) pre-coated ELISPOT plates and stimulated with the GalHyK264 peptide for 24 h. Bound cytokines were detected by biotinylated anti-IFNγ (R46-A2) (Mabtech, #3321-6-1000) or anti-IL17A (TC11-8H4) (Biolegend, #507002) or anti-IL-4 (BVD6-24G2) (BD, #1244-360) and measured by dissociation-enhanced time-resolved fluorometry (excitation 360/40 and emission 620/40).

5.7. Antigen presentation assays

Splenocytes were co-cultured with COL2-specific T cell hybridoma (HCQ.3 or HCQ.4 hybridoma) cells for 24 h in the presence of heat-denatured COL2 (using HCQ.3), K264 (using HCQ.4) or GalHyK264 peptide (using HCQ.3). IL2 production in the supernatant was detected by ELISA. Briefly, culture supernatant was subjected to anti-IL2 (Jes6-1A2, homemade, 1 μg/ml) pre-coated plates, and captured IL2 was detected by biotinylated anti-IL2 (Jes6-5H4, homemade, 1 μg/ml) with Eu-labeled streptavidin (PerkinElmer, #1244-360) and measured by dissociation-enhanced time-resolved fluorometry (excitation 360/40 and emission 620/40, Synergy-2; BioTek Instruments). Recombinant
murine IL2 (Peprotech, #210-12) was used as standard.

5.8. B cell isolation and in vitro differentiation

B cells were isolated by magnetic-activated cell sorting (MACS) techniques following the protocol provided by Miltenyi Biotec (Order no.130-121-30, CD19 MicroBeads, mouse). Briefly, splenocytes were resuspended in MACS buffer and incubated with CD19 MicroBeads for 10 min, then subjected to pre-lensed column (Miltenyi Biotec, #130-042-401). After washing with MACS buffer for 3 times, B cells were eluted by flushing 5 ml of MACS buffer using the plunger, then resuspended in complete RPMI supplemented with 10% FCS and penicillin/streptomycin.

For lipopolysaccharide (LPS) stimulation, B cells were cultured in TC-treated 96 well U-bottom plates (Falcon®, #353077) in complete RPMI media further supplemented with 10 μg/ml LPS (Sigma-Aldrich, #L2880) for 64 h. 10 μM hydrogen peroxide (H₂O₂, Sigma-Aldrich, #H3410) and 10 μM GS (GSK2795039, Sigma-Aldrich, #SML2770) were added in after 40 h incubation. For CD40L stimulation, B cells were cultured 84 h in CD40L expressing irradiated (50 Gy) fibroblasts (kind gift from Ritu Carsetti, B Cell Pathophysiology Unit, Immunology Research Area, Bambino Gesù Children’s Hospital IRCCS, 00146 Rome, Italy) pre-coated TC-treated 96 well U-bottom plates, and further supplemented with 50 ng/ml murine IL21 (Peprotech, #210-21) with or without 50 ng/ml murine IL4 (Peprotech, #214-14) with or without 5 μg/ml goat anti-mouse IgM (Fab)² (anti-μ) (Jackson Immuno Research, #115-006-075) as indicated. Cells were cultured at 37 °C in humidified air with 5% CO₂. Culture supernatants were aspirated carefully and checked for total IgM antibodies. Cells were collected for flow cytometric analysis.

5.9. B cell transfer experiment

B cells were MACS-sorted from spleens of male B6N.CD45 mice, male B10Q.CD45 Ncf4−/−/− and male B10Q.CD45 Ncf4+/-/+/- mice then adjusted to the same concentration in PBS (Thermo fisher, #14190169). For chimeric B cell transfer experiment (performed in Fig. 5A–G), CD45.1* B cells (from B6N.CD45 mice) and CD45.2* B cells (from B10Q.CD45 Ncf4−/−/− or B10Q.CD45 Ncf4+/-/+/- mice) were equally mixed and transferred to recipient B10Q.CD45 Ncf4−/−/− mice intravenously. For experiments performed in Fig. 5H, B cells were not mixed and transferred to recipient B10Q.CD45 Ncf4−/−/− mice separately. A total of 10 million B cells were transferred to each recipient mouse. B cell viability and purity after sorting were checked by flow cytometry. Blood from recipient mice was taken for checking the transfer efficacy. 2 days after transfer, mice were induced CIA following the protocol described above (without boost).

5.10. Statistical analysis

Experiments pooled had balanced groups and no data has been excluded. Data were expressed as mean ± SEM and analyzed using GraphPad Prism (Version 8.0). Mann-Whitney test was employed to analyze significant differences between two groups. Kruskal-Wallis test and Dunn’s multiple comparisons test were used for comparing differences between three groups. Pearson correlation analysis was used to analyze correlation between antibody titers and disease severity. Multiple Student’s t tests with Holm-Sidak’s comparison correction were used to determine the significant differences in the disease severity and incidence of CIA on different DPs. P-values smaller than the significance level (set to 0.05) are indicated by asterisks (*p<0.05, **p<0.01, ***p<0.001, and ****p<0.0001).

Author contributions

CH designed the research, performed most of the experiments, including acquiring and analyzing data. HL contributed with Pil related experiments and part of in vitro experiments. AC, ML and QL contributed with part of long-term CIA experiments. AK provided the irradiated CD40L expressing fibroblasts and contributed with designing the in vitro experiment. JX, SM and 2Y contributed with experiments design and manuscript revision. RH designed the research, analyzed the data, revised the manuscript, supervised, and takes the overall responsibility of the study. All authors revised and approved the manuscript.

Declaration of competing interest

The authors declare no competing financial interests.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.redox.2022.102422.

References

[1] M.C. Dinauer, S.H. Orkin, R. Brown, A.J. Jesaitis, C.A. Parkos, The glycoprotein encoded by the X-linked chronic granulomatous disease locus is a component of the neutrophil cytochrome b complex, Nature 327 (6124) (1987) 717–720.
[2] O. Sarrela, T. Kelkka, A. Pizzolla, M. Hultqvist, R. Holmdahl, NOX2 complex-derived ROS as immune regulators, Antioxidants Redox Signal. 15 (8) (2011) 2197–2204.
[3] T. Ueyama, T. Tsutsumo, T. Kasawaki, S. Tsujibe, Y. Shirai, H. Sumimoto, T.L. Leto, N. Saito, A regulated adaptor function of p40phox: distinct p67phox membrane targeting by p40phox and by p7phox, Mol. Biol. Cell 18 (2) (2007) 441–454.
[4] R.V. Shahelin, A. Iuriin, K.S. Bruzik, D. Murray, W. Cho, Membrane binding mechanisms of the PX domains of NADPH oxidase p40phox and p47phox, J. Biol. Chem. 278 (16) (2003) 14469–14479.
[5] P. Stampoulis, T. Ueda, M. Matsutomo, H. Terasawa, K. Miyano, H. Sumimoto, L. Shimada, Atypical membrane-embedded phosphatidylinositol 3,4-bisphosphate PI(3,4)P2-binding site on p47(phox) Phox homology (PX) domain revealed by NMR, J. Biol. Chem. 287 (21) (2012) 17848–17859.
[6] E.M. Song, L. Bouchal, E. Hudik, R. Le Bars, O. Nusse, S. Dupre-Crochet, Phosphoinositol 3-phosphate acts as a timer for reactive oxygen species production in the phagosome, J. Leukoc. Biol. 101 (5) (2017) 1155–1168.
[7] J.D. Pollock, D.A. Williams, M.A. Gifford, L.L. Li, X. Du, J. Fisherman, S.H. Orkin, C.M. Doerschuk, M.C. Dinauer, Mouse model of X-linked chronic granulomatous disease, an inherited defect in phagocyte superoxide production, Nat. Genet. 20 (2) (1995) 202–209.
[8] A. van de Geer, A. Nieto-Plantat, D.B. Kuhns, A.T. Tool, A.A. Arias, M. Bouaziz, M. de Boer, J.L. Franco, R.P. Gazendam, J.L. van Hamme, M. van Houdt, K. van Leuven, P.J. Verkuiljen, T.K. van den Berg, J.F. Aflate, C.A. Arango-Franco, V. Batura, A.R. Bernasco, B. Boardman, C. Booth, S.O. Burns, F. Cabarcas, N. Cesana, F. Charbit-Henrion, A. Corveleyn, C. Deswarte, M.E. Azcoiti, D. Foell, I. Shimada, Atypical membrane-embedded phosphatidylinositol 3,4-bisphosphate PI(3,4)P2-binding site on p47(phox) Phox homology (PX) domain revealed by NMR, J. Biol. Chem. 287 (21) (2012) 17848–17859.
C. He et al.

[27] E.J. Kunkel, E.C. Butcher, Plasma-cell homing, Nat. Rev. Immunol. 3 (10) (2003) 1431–1437.

[26] M. Bertolotti, S.H. Yim, J.M. Garcia-Manteiga, S. Masciarelli, Y.J. Kim, M.H. Kang, J. Bagaitkar, E.A. Barbu, L.J. Perez-Zapata, A. Austin, G. Huang, S. Pallat, M. Helminen, L. Kainulainen, P. Olofsson, S. Jalkanen, R. Lahesmaa, M.M. Souto-Carneiro, R. Holmdahl, Reactive oxygen species deficiency induces autoimmunity with type I interferon signature, Antioxidants Redox Signal. 21 (16) (2014) 2233–2245.

[20] J. Zhao, J. Ma, Y. Deng, J.A. Kelly, K. Kim, S.Y. Bang, H.S. Lee, Q.Z. Li, E. K. Wakens, R. Qiu, M. Liu, J. Guo, Z. Li, W. Tan, A. Rasmussen, C.J. Lessard, K. L. Stilv, B.H. Hahn, J.M. Grossman, D.L. Kamen, G.S. Gilkeson, S.C. Bae, P. M. Gaffney, N. Shen, B.P. Tao, A. Misra, E. B. Mortensen, M. Hultqvist, J. Holmberg, P. Olofsson, R. Holmdahl, T cell surface redox levels determine T cell reactivity and arthritis susceptibility, Proc. Natl. Acad. Sci. U. S. A. 103 (34) (2006) 12831–12836.

[19] L.V. Crozet, J.D. Mate, A.A. Arias, H. Zhao, L.A. Quilliam, M.C. Dinauer, J. S. Blum, Cutting edge: NADPH oxidase modulates MHC class II antigen presentation by B cells, J. Immunol. 189 (8) (2012) 3800–3804.

[18] R.A. Ehner, M.J. Shlomchik, Germinal center and extrafollicular B cell responses in vaccination, immunity, and autoimmunity, Immunity 53 (6) (2020) 1136–1150.

[17] C.O. Jacob, M. Eisenstein, M.C. Dinauer, W. Ming, J. Yu, J. Ma, Y. Deng, J.A. Kelly, K. Kim, S.Y. Bang, H.S. Lee, Q.Z. Li, E. K. Wakens, R. Qiu, M. Liu, J. Guo, Z. Li, W. Tan, A. Rasmussen, C.J. Lessard, K. L. Stilv, B.H. Hahn, J.M. Grossman, D.L. Kamen, G.S. Gilkeson, S.C. Bae, P. M. Gaffney, N. Shen, B.P. Tao, A. Misra, E. B. Mortensen, M. Hultqvist, J. Holmberg, P. Olofsson, R. Holmdahl, T cell surface redox levels determine T cell reactivity and arthritis susceptibility, Proc. Natl. Acad. Sci. U. S. A. 103 (34) (2006) 12831–12836.

[16] C.P. Marques, P. Kap, D.R. Hutton, C. Hindinger, S.L. Nutt, R.M. Ransohoff, T. W. Phares, S.A. Stohlmann, C.G. Bergmann, CXCR3-dependent plasma blast migration to the central nervous system during viral encephalomyelitis, J. Virol. 85 (13) (2011) 6136–6147.

[15] L.M. Olsson, A.K. Lindqvist, H. Kallberg, L. Padyukov, H. Burkhardt, L. Alfredsson, J. Holmberg, J. Tordsson, S. Lu, B. Åkerström, R. Holmdahl, Positional identification of Ncf1 as a gene that regulates arthritis severity in rats, Nat. Genet. 33 (1) (2003) 25–32.

[14] L. Sivils, B.H. Hahn, J.M. Grossman, D.L. Kamen, G.S. Gilkeson, S.C. Bae, F. Ho, J.E. Lortan, I.C. MacLennan, D.L. Armstrong, R. Zidovetzki, Lupus-susceptible NZM2410 mice. I. Different regulation of autoreactive vs. non-autoreactive anti-type II collagen T cells in the DBA/1 mouse, Eur. J. Immunol. 20 (5) (1990) 1061–1073.

[13] J. Britton, O. Kallberg, L. Alfredsson, J. Holmberg, J. Tordsson, S. Lu, B. Åkerström, R. Holmdahl, Positional localization of the Ncf1 gene controlling oxidative burst response and arthritis severity in rats, Antioxidants Redox Signal. 14 (12) (2011) 2373–2383.

[12] J. Kunkel, E.J. Kunkel, E.C. Butcher, Plasma-cell homing, Nat. Rev. Immunol. 3 (10) (2003) 1431–1437.

[11] L.M. Olsson, A. Nerstedt, A.K. Lindqvist, S.C. Johansson, P. Medstrand, P. Olofsson, R. Holmdahl, Copy number variation of the gene NCF1 is associated with rheumatoid arthritis, Antioxidants Redox Signal. 16 (1) (2012) 71–78.

[10] D. Kienhofer, M. Hoffmann, M. Linja, K. Wing, O. Sareila, M. Hultqvist, J. Holmberg, P. Olofsson, R. Holmdahl, Lack of reactive oxygen species breaks T cell tolerance to collagen type II and allows development of arthritis in mice, J. Immunol. 179 (3) (2007) 1431–1437.

[9] K.A. Gelderman, M. Hultqvist, J. Holmberg, P. Olofsson, R. Holmdahl, T cell surface redox levels determine T cell reactivity and arthritis susceptibility, Proc. Natl. Acad. Sci. U. S. A. 103 (34) (2006) 12831–12836.

[8] F. Ho, J.E. Lortan, I.C. MacLennan, M. Khan, Positional cloning of the Ncf1 gene controlling oxidative burst response and arthritis severity in rats, Antioxidants Redox Signal. 7 (5) (2007) 896–906.

[7] T.P. Zhang, R. Li, Q. Huang, H.F. Pan, D.Q. Ye, X.M. Li, Association of NC2F, NCF4, and CYBA gene polymorphisms with rheumatoid arthritis in a Chinese population, J Immunol Res (2020), 8528976, 2020.

[6] P. Olofsson, J. Holmberg, J. Tordsson, S. Lu, B. Åkerström, R. Holmdahl, Positional identification of Ncf1 as a gene that regulates arthritis severity in rats, Nat. Genet. 33 (1) (2003) 25–32.

[5] M. Hultqvist, O. Sareila, F. Vilhardt, U. Norin, L.M. Olsson, P. Olofsson, U. Hellman, R. Holmdahl, Positioning of a polymorphic quantitative trait nucleotide in the Ncf1 gene controlling oxidative burst response and arthritis severity in rats, Antioxidants Redox Signal. 14 (12) (2011) 2373–2383.

[4] T. Kericka, D. Kienhofer, M. Hoffmann, M. Linja, K. Wing, O. Sareila, M. Hultqvist, E. Lajalda, Z. Chen, J. Vasconcelos, E. Neves, M. Guedes, I. Marques, G. Konouk, M. Helminen, L. Kainulainen, P. Olofsson, S. Falken, R. Lahesmaa, M.M. Souto-Carneiro, R. Holmdahl, Reactive oxygen species deficiency induces autoimmunity with type I interferon signature, Antioxidants Redox Signal. 21 (16) (2014) 2233–2245.

[3] C.O. Jacob, M. Eisenstein, M.C. Dinauer, W. Ming, J. Yu, S. John, F. Quismorio Jr., A. Reiff, B.L. Myones, J. Tordsson, R. Holmdahl, Positioning of a polymorphic quantitative trait nucleotide in the Ncf1 gene controlling oxidative burst response and arthritis severity in rats, Antioxidants Redox Signal. 14 (12) (2011) 2373–2383.

[2] J. Britton, O. Kallberg, L. Alfredsson, J. Holmberg, J. Tordsson, S. Lu, B. Åkerström, R. Holmdahl, Positional identification of Ncf1 as a gene that regulates arthritis severity in rats, Nat. Genet. 33 (1) (2003) 25–32.

[1] L.M. Olsson, A. Nerstedt, A.K. Lindqvist, S.C. Johansson, P. Medstrand, P. Olofsson, R. Holmdahl, Copy number variation of the gene NCF1 is associated with rheumatoid arthritis, Antioxidants Redox Signal. 16 (1) (2012) 71–78.