Immune signatures of prodromal multiple sclerosis in monozygotic twins

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The tremendous heterogeneity of the human population presents a major obstacle in understanding how autoimmune diseases like multiple sclerosis (MS) contribute to variations in human peripheral immune signatures. To minimize heterogeneity, we made use of a unique cohort of 43 monozygotic twin pairs clinically discordant for MS and searched for disease-related peripheral immune signatures in a systems biology approach covering a broad range of adaptive and innate immune populations on the protein level. Despite disease discordance, the immune signatures of MS-affected and unaffected cotwins were remarkably similar. Twinship alone contributed 56% of the immune variation, whereas MS explained 1 to 2% of the immune variance. Notably, distinct traits in CD4+ effector T cell subsets emerged when we focused on a subgroup of twins with signs of subclinical, prodromal MS in the clinically healthy cotwin. Some of these early-disease immune traits were confirmed in a second independent cohort of untreated early relapsing-remitting MS patients. Early involvement of effector T cell subsets thus points to a key role of T cells in MS disease initiation.

multiple sclerosis | monozygotic twins | immune phenotyping | autoimmunity | biomarker

Research into human autoimmunity is facing a number of particular challenges. First, in contrast to inbred animal strains housed under controlled conditions, the human population is exceedingly heterogeneous, with a high degree of genetic and environmental diversity. Second, human autoimmune diseases are usually diagnosed long after initiation of the disease process, so it is impossible to differentiate between early (primary) and late (secondary) immune processes. Third, human studies mostly rely on the blood as the main source of immune cells because access to human tissues is usually limited. Last but not least, ethical constraints limit the scope of functional (“mechanistic”) human studies compared to sophisticated animal systems. For these and other reasons it has been very difficult to capture the early footprints (“signatures”) of nascent human autoimmune reactions in the blood, the most accessible peripheral immune compartment. This holds particularly true for organ-specific, “compartamentalized” autoimmune processes, such as autoimmune diabetes, rheumatoid arthritis, or multiple sclerosis (MS).

MS is a prominent example of such a compartmentalized autoimmune disease as it affects the central nervous system (CNS), which is partly secluded from the peripheral immune system by the blood–brain barrier (1–3). Classic experiments in animals showed that experimental autoimmune encephalomyelitis (EAE), the most widely used model of MS, can be induced in naïve recipient animals by injecting purified myelin autoantigen-specific CD4+ effector T cells (4, 5). However, consistent with the excessive heterogeneity of the human population, it has so far not been possible to identify any unifying “target autoantigen” in human MS. There are additional notable differences between EAE models and human MS, including the well-known preponderance of clonally expanded CD8+, rather than CD4+, T cells in MS brain lesions, and the presence of oligoclonal immunoglobulins in the cerebrospinal fluid (CSF) of MS patients (6–9). Yet despite knowledge about distinct MS-associated features in affected tissues, identification of an MS-associated peripheral immune signature has remained elusive.

The MS Twin Study included organization, recruitment, and patient care; A.F.-H., B.E.-W., T.K., and H.W. were coinvestigators of the MS Twin Study; S.G.M., H.W., and L.K. characterized MS patients (6–9). Classic experiments in animals showed that experimental autoimmune encephalomyelitis (EAE), the most widely used model of MS, can be induced in naïve recipient animals by injecting purified myelin autoantigen-specific CD4+ effector T cells (4, 5). However, consistent with the excessive heterogeneity of the human population, it has so far not been possible to identify any unifying “target autoantigen” in human MS. There are additional notable differences between EAE models and human MS, including the well-known preponderance of clonally expanded CD8+, rather than CD4+, T cells in MS brain lesions, and the presence of oligoclonal immunoglobulins in the cerebrospinal fluid (CSF) of MS patients (6–9). Yet despite knowledge about distinct MS-associated features in affected tissues, identification of an MS-associated peripheral immune signature has remained elusive.

Apart from the heterogeneity of the human population, built-in delays in the diagnostic process of autoimmune diseases have hindered the identification of peripheral disease-specific signatures. In MS for example, diagnosis strictly requires the presence of clinical symptoms according to prevailing consensus criteria (10). However, the appearance of symptoms can be preceded by...
a long-lasting prodromal phase of clinically silent neuroinflammation. This is detected only very rarely and usually only by chance when unexpected silent brain lesions are observed on MRI performed for an unrelated medical reason, a setting referred to as “radiologically isolated syndrome” (11).

In searching for peripheral MS-associated immune signatures, we here took advantage of a unique cohort of 43 monozygotic twin pairs who are discordant for MS. That is, one cotwin in each pair carries the diagnosis of MS, whereas the other cotwin is clinically healthy. We reasoned that pairwise comparisons of these monozygotic twins would eliminate genetic heterogeneity for each pair, and in addition reduce environmental heterogeneity as the twins included in this study were raised in the same household. In a first step of analysis, we found that the immune signatures of MS-affected and unaffected cotwins were remarkably similar, whereas age, sex, and MS disease status together contributed less than 10% of the variation of immune traits. In a second step, we leveraged another distinct advantage of the MS-discordant twin cohort: The healthy cotwins of MS-affected twins have a maximum familial risk of developing MS and therefore an increased probability of showing signs of prodromal MS (12). Indeed, several clinically unaffected cotwins showed distinct signs of early subclinical neuroinflammation (SCNI) on MRI. By focusing on these subjects with early neuroinflammation, we were able to detect an early influence of MS on the peripheral immune signature, using high-resolution flow cytometry-based immune phenotyping followed by unbiased machine-based analysis and dimensionality reduction algorithms.

**Results**

**High Degree of Similarity in Peripheral Immune Signatures within Twin Pairs Despite Disease Discordance.** In this study, we evaluated MS-related changes in peripheral immune signatures in a cohort of 43 monozygotic twin pairs clinically discordant for MS (Fig. 1A), as a twin approach offers the important advantage to control for genetic as well as intrinsic factors as age and sex, as well as early environmental factors. All 43 monozygotic twin pairs were of Caucasian origin and were raised together until adulthood and shared a variety of early environmental factors. Demographic data, as well as information with regard to MS disease course and treatment, are depicted in Table 1. We took advantage of a comprehensive deep functional immune-cell phenotyping platform by using multicolor flow cytometry on frozen peripheral blood mononuclear cell (PBMC) specimens. In this system immunology approach, 13 9- to 10-color immunophenotyping panels were used for the characterization of major immune cell subpopulations and their interactions. The characterization was performed using high-resolution flow cytometry-based immune phenotyping followed by unbiased machine-based analysis and dimensionality reduction algorithms.

**Fig. 1.** In-depth characterization of monozygotic twins discordant for MS. (A) Twin study set-up of monozygotic twin pairs (n = 43) discordant for MS: Healthy (TWINHD, gray) and MS (TWINMS, red) twin. Sample processing; cryoconserved PBMC of study participants were processed via flow cytometry in 13 panels of 10 colors each. For unbiased nonlinear dimensionality reduction via bh-SNE, LMDs were transformed, normalized, and merged per panel and randomly subsampled in viSNE, an implementation in Matlab. For visual representation of single-cell data and identification of population clusters, a Phenograph algorithm was used; fold-change was calculated per cluster comparing compiled data from all TWINHD to TWINMS; a regulation greater than twofold (up-regulation in red, down-regulation in green) is visualized in a heatmap. (B) Hierarchical illustration of the defined populations (n = 141) analyzed within this study. Red dots illustrate the respective main populations within the adaptive and innate compartment, further subpopulations are indicated by blue dots.
maturation and functional status (13, 14). Longitudinal repeated measurements (n = 26) of PBMCs derived from a study-independent healthy individual during the total acquisition period of 4 wk (mean variability of traits r = 0.05) revealed robustness of data acquisition (Dataset S4). To accommodate for sample-to-sample acquisition variability, we used a linear normalization algorithm and applied it on a single-file level. The normalized single-cell data were then visualized by Barnes–Hut stochastic neighbor embedding (bh-SNE) and Phenograph, a machine-based algorithm for clustering and dimensionality reduction of high-dimensional single-cell data (15, 16). Fig. 1B illustrates the immune cell parameters as determined by conventional gating, finally encompassing 141 different immune phenotypes, which we here refer to as immune cell traits (17).

To detect and visualize distinct immune signatures in healthy twins (TWINHD) (Fig. 2A, second row) and MS-affected (TWINMS) (Fig. 2A, second row) cotwins, we used unsupervised clustering to identify major immune cell populations within each panel (Fig. 2A) by the Phenograph algorithm. The algorithm identified 30 distinct immune cell clusters (as described in Materials and Methods) in high-dimensional space based on compiled TWINHD and TWINMS single-cell data. We did not observe any differences between TWINHD and TWINMS in any of the immune cell traits encompassed by this analysis as shown in the heatmaps (Fig. 2A, third row). This was corroborated by conventional manual gating approaches yielding the frequencies of the main immune cell populations and subsequent Pearson correlation analysis (Fig. 2A and SI Appendix, Fig. S1A). Notably, also pairwise comparison within individual twin pairs revealed a remarkable similarity in the immune traits for all major immune cell populations, with a slight tendency for differential patterns only found in the B cell subset of treated MS cotwins (Fig. 2B).

We next analyzed immune cell lineages in more detail, by considering all (n = 13) (Fig. 2C) immunophenotyping panels together. Across all flow cytometry panels, Phenograph identified 352 discrete immune cell clusters in high-dimensional space, which we used to reveal potential differences between TWINHD and TWINMS immune cell signatures with higher resolution. Still, we did not observe any distinctly emerging changes (i.e., defined as more than twofold) per cluster between TWINHD and TWINMS, again pointing toward a strong similarity within each twin pair (Fig. 2 C and D).

When focusing our analysis on those twin pairs without any MS treatment—to exclude potential treatment-related confounding factors—we again did not detect any relevant changes within each cluster comparing TWINHD to TWINMS groups (SI Appendix, Fig. S1B).

We also performed functional analyses of major subsets, using intracellular cytokine expression analysis following short-term stimulation. We found slightly lower frequencies of cytokine-producing CD4+ and CD8+ T memory subsets in the MS cotwins; however, this difference was completely abolished when focusing on those MS cotwins not receiving any immune therapy at the time of blood sampling, hence suggesting that these effects were most likely treatment related (SI Appendix, Fig. S1C).

**Twinship Explains the Majority of the Variance in Peripheral Immune Traits.** We next aimed to decipher and quantify the distinct individual effects of defined factors, such as twinship, MS status, age, gender, and cytomegalovirus (CMV) serostatus on the variance of all innate and adaptive immune cell traits investigated using a linear mixed-model approach (Fig. 3A). The factor “twinship” encompasses both the genetic background, and shared local environmental factors, such as childhood infections or diet, that play a major role in early development of immunity (18, 19). Importantly, twinship explained the vast majority of the variance for each of the 141 investigated immune-cell traits, its influence ranging from 11 to 89% of the explained variance (Fig. 3B). In contrast, age, sex, CMV, and MS status taken together had a minor contribution (11%) to the variance of each parameter (Fig. 3 B and C and SI Appendix, Fig. S2). Overall, twinship was responsible on average for the major proportion (56%) of the variance, whereas the MS status explained 1%, age 4%, CMV status 3%, and sex 2% of the proportion of the variance of immune traits (Fig. 3D). To search for subtle differences in the contribution of each factor, all parameters were grouped according to their immunological “ancestry” (Fig. 1B). Again, the MS status alone explained minor proportions of the variation, ranging from 1% in conventional T cells and innate immune cells, to 2% in B cells, monocytes and dendritic cells (DC) (Fig. 3 C and D). Interestingly, the contribution of twinship to the variance of B cell traits was slightly less pronounced (Fig. 3 C and D), albeit

**Table 1. Basic clinical characteristics of patient cohorts**

<table>
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<tr>
<th>Number</th>
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<th>10</th>
<th>55</th>
<th>60</th>
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<td>44 (20–67)</td>
<td>32.6 (18–56)</td>
<td>33.1 (18–55)</td>
<td>33.3 (21–56)</td>
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<td></td>
<td></td>
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<td>0</td>
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<tr>
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<td>14 y (1–30)</td>
<td>1.6 mo (0–10)</td>
<td>2.7 mo (0–10)</td>
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<td>1.3 (0–2.5)</td>
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<td></td>
<td></td>
</tr>
<tr>
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<td>5</td>
<td>55</td>
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NA, not applicable; PPMS, primary progressive MS; SPMS, secondary progressive MS.

*Tested pairs n = 36.
†Tested pairs n = 42.

When focusing our analysis on those twin pairs without any MS treatment—to exclude potential treatment-related confounding factors—we again did not detect any relevant changes within each cluster comparing TWINHD to TWINMS groups (SI Appendix, Fig. S1B).
Fig. 2. Frequencies of immune cell populations and functional properties in monozygotic twins discordant for MS. (A) PhenoGraph-maps compare single cell clusters of major immune cell populations of compiled TWINHD (row 1) to TWINMS (row 2) data indicated by black circles (basic panel n = 43 pairs; ILC panel n = 34 pairs). Heatmap (row 3) illustrates the fold-change per cluster between TWINMS and TWINHD, >twofold up-regulation (red) and >twofold down-regulation (green) would be indicated within the black circles. Corresponding conventional flow cytometry data comparing TWINMS to TWINHD are depicted in row 4. (B) Illustration of pairwise differences within individual twin pairs for all major immune populations based on conventional flow cytometry data; heatmap illustrates the difference in percentage. Immune therapy of MS twins (if applicable) is indicated below, IFN-β, natalizumab (NAT), and glatiramer acetate (GLAT), or no treatment (untreated). (C) Unbiased single-cell analysis of flow cytometry data via viSNE for each individual panel (panels = 13); PhenoGraph algorithm identified in total 352 clusters of compiled data from TWINMS and TWINHD groups (basic-, Th1-, Th17-, stimulation 2 panel n = 43; B cell panel n = 41 pairs; CD8+ panel n = 39 pairs; DC panel n = 30 pairs; ILC panel n = 34 pairs; monocyte panel 1+2 n = 37 pairs; stimulation 1 panel n = 31 pairs; Treg panel n = 42 pairs); illustrations display the frequency per cluster within each panel. (D) Corresponding heatmap illustrates the fold-change between TWINMS and TWINHD groups for each individual cluster; twofold change line (dotted red line), no change line (dotted gray line) are indicated, Graph values represent mean ± SD. Statistical significance was evaluated by Wilcoxon matched-pairs signed rank test; n.s.: not significant.
Fig. 3. Effects of age, sex, CMV, MS status, and twinship on the variation of immune cell traits in monozygotic twins. (A) Illustration of our approach of quantification of the proportion of variance explained ($R^2$) by fixed factors age, sex, CMV, and MS defined as the marginal $R^2$ and additional incorporation of twinship as random effect (conditional $R^2$) in a linear mixed model. (B) Dot plot graphs demonstrate the marginal versus conditional $R^2$ for each immune cell trait in TWINMS and TWINHD cohorts ($n = 43$ pairs). (C) Dot plot graphs demonstrate the marginal versus conditional $R^2$ for each individual immune trait within the respective major immune cell population. (D) Bar graphs illustrate the mean proportions of variance ($R^2$) explained by the different factors age, sex, CMV, and twinship for all major immune cell populations; residual unexplained variance is displayed in violet.
We observed a significant ICC rise in the CD4+ but not CD8+ compartment of the T-cell subset in the CD4+ effector subsets as described in intrathecal IgG synthesis (8). A significant proportion of clinically unaffected cotwins (n = 10) presented signs of SCNI, suggesting the presence of prodromal MS. A group of clinically healthy cotwins (n = 14) without any signs of SCNI were used as controls for further analysis (Fig. 4A). We first repeated our unbiased cluster analysis of healthy twins versus MS twins by only incorporating datasets from healthy twins without any signs of SCNI; however, this did not result in the identification of differentially regulated clusters (SI Appendix, Fig. S3). We then hypothesized that if any immune cell trait is indeed influenced by MS, the extent of its correlation would be higher within SCNI-MS twin pairs than in healthy donors (HD)-MS twin pairs based on the premise that SCNI represents the earliest detectable stage of MS. We therefore performed a Spearman correlation analysis adjusted for MS status, comparing the correlation coefficient of SCNI-MS twin pairs with the correlation coefficient of HD-MS twin pairs for all immune cell subsets (n = 141) (SI Appendix, Fig. S4A). As hypothesized, we observed a significant increase in the degree of correlation in SCNI-MS twin pairs particularly within the adaptive compartment (i.e., all T cells and, more specifically, in CD4+ and CD8+ T cell subsets). The degree of correlation was also slightly increased for the monocyte/dendritic cell subset.

To further corroborate these findings by another statistical approach, we calculated the intraclass correlation coefficient (ICC) (Fig. 4B) for each immune cell trait based on a linear mixed-model adjusted for MS, and corrected the comparison between SCNI-MS and HD-MS twin cohorts for the hierarchical dependency of parameters (SI Appendix, Fig. S4B). The advantage of this approach is that the mixed model was additionally adjusted for the interdependence of immune parameters as described in Materials and Methods. Comparison of the ICC of the major immune cell populations revealed a significant increase in the SCNI-MS twin pairs compared to the HD-MS twin pairs for T cells but neither for B cells nor for the major innate cell populations (Fig. 4B and SI Appendix, Fig. S4C), again supporting the concept that the degree of correlation is enhanced in SCNI-MS twin pairs as compared to HD-MS twin pairs. Within the T cell compartment, we observed a significant ICC rise in the CD4+ but not CD8+ T cell subsets (Fig. 4B and SI Appendix, Fig. S5A).

For further corroboration of this concept we made use of unbiased t-distributed stochastic neighbor embedding (t-SNE) dimensionality reduction of our flow cytometry data to calculate the biological distance between each individual twin pair as an indicator of the degree of similarity within predefined CD4+ T cell subsets as described in Materials and Methods (a representative example of such a calculation is depicted in Fig. 4C and SI Appendix, Fig. S5B). Indeed, we observed a significant difference in the biological distance between SCNI-MS twin pairs and HD-MS twin pairs for CD4+ effector subsets (Fig. 4C), again pointing toward a higher degree of similarity within the SCNI-MS twin pairs. Heatmap visualization of pairwise differences (in percent) for individual parameters within the CD4+ effector subsets further illustrated the increased degree of similarity within the SCNI-MS twin pairs, which was particularly pronounced within proinflammatory CD4+ T-cell subsets, such as Th17 and Th1 cell populations (Fig. 4D).

Validation of MS-Related Immune Traits in an Early MS Replication Cohort. Finally, we wanted to determine whether our approach of analyzing individuals with prodromal MS might facilitate identification of potentially disease-associated peripheral immune signatures in a replication cohort of subjects with either clinically isolated syndrome (CIS) or early clinically manifest relapsing-remitting MS (RRMS) according to the 2010 Mc Donald criteria (21), that were all treatment-naïve at the time of blood sampling (Fig. 4E). In this cohort, biosample acquisition took place within the first 3 years from the time of clinically defined disease onset. We were able to obtain flow cytometry data of our key candidate populations identified in Fig. 4D, in 55 patients with clinically isolated syndrome and 60 treatment-naive early RRMS patients, as well as 71 age- and sex-matched healthy controls (Table 1). Indeed, conventional flow cytometry analysis revealed significant MS-related changes in some of the previously identified immune traits from our twin approach, such as Th1 cells, migratory Th17 cells (MCAM+), effector Th17 cells, migratory Th1 cells (CD195+), as well as Th17.1 cells (CD195+), as well as Th17.1 cells (CD195+) (Fig. 4E). To further corroborate these findings by another statistical approach, we calculated the intraclass correlation coefficient (ICC) (Fig. 4F) for each immune cell trait based on a linear mixed-model adjusted for MS, and corrected the comparison between SCNI-MS and HD-MS twin cohorts for the hierarchical dependency of parameters (SI Appendix, Fig. S4B). The advantage of this approach is that the mixed model was additionally adjusted for the interdependence of immune parameters as described in Materials and Methods. Comparison of the ICC of the major immune cell populations revealed a significant increase in the SCNI-MS twin pairs compared to the HD-MS twin pairs for T cells but neither for B cells nor for the major innate cell populations (Fig. 4B and SI Appendix, Fig. S4C), again supporting the concept that the degree of correlation is enhanced in SCNI-MS twin pairs as compared to HD-MS twin pairs. Within the T cell compartment, we observed a significant ICC rise in the CD4+ but not CD8+ T cell subsets (Fig. 4B and SI Appendix, Fig. S5A).

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Discussion

In our study we aimed at capturing autoimmunity-related changes in the peripheral immune system in a systems biology approach. To this end, we took advantage of a MS-discordant monozygotic twin cohort, which allows for controlling genetic as well as intrinsic factors, including shared early environmental influences on immune cell traits. At the level of the overall cohort, immune signatures of MS-affected and unaffected twins were remarkably similar despite discordance for an autoimmune disease. However, when we focused on a subgroup of clinically healthy cotwins showing signs of early subclinical neuroinflammation, a distinct MS-related immune signature emerged.

Dominance of Twinship Over MS Disease Status. Using a comprehensive flow cytometry-based immune phenotyping approach and linear mixed models, we determined the relative impact of age, sex, CMV status, MS status, and twinship on the variation of each immune cell trait. Our analysis revealed that the impact of twinship was dominant, ranging from 11% to a maximum 89%, with a mean of 56% of explained variance for all immune traits investigated. Other known factors—such as age, sex, and CMV status—had an influence on average of 4% for age, 2% for sex, and 3% for CMV, thus confirming their acknowledged effects on cellular immune compartments as, for example, recent thymic emigrant CD4+ T cells for age (18, 22). In comparison, MS exerted an average influence of 1%, indicating a similar degree of impact as compared to the established factors age, sex, and CMV status.

A distinct influence of twinship on the variation of immune cell traits was expected based on studies in healthy individuals, especially in light of the known impact of genetic as well as local environmental factors on the immune system (18, 19, 23–26). Along this line, several recent studies provided evidence of a strong genetic influence on immune cell traits: One study revealed that adaptive immune traits—in particular CD4+ and CD8+ T cells—were more influenced by genetics than innate traits (23). Another study also revealed a substantial impact of genetic factors on immune traits in a cohort of 1,620 Sardinians (27). Moreover, an analysis from the human functional genomics project demonstrated that within the lymphocyte population, genetics explained a higher percentage of variation for T cells (i.e., 30%) than for
Fig. 4. SCNI analysis identifies effector CD4 populations as differentially regulated populations in the initiation of MS. (A) Illustration of cohort division into SCNI-MS twin pairs (n = 10, beige) and HD-MS twin pairs (n = 14, gray) as described in Materials and Methods. (B) Calculation of the ICC for each immune trait in SCNI-MS twin pairs (n = 10, beige) and HD-MS twin pairs (n = 14, gray) based on linear mixed model adjusted for MS. To account for hierarchical dependencies between immune cell populations, analyses incorporate corrections using correlation matrices per sub collective, as described in Materials and Methods. (C) Representative example of t-SNE dimensionality reduction of conventionally analyzed flow cytometry populations comparing the biological distance (gray line) between HD-cotwins (gray rectangle, n = 10) and SCNI cotwins (beige circle, n = 9) to their respective MS cotwin (red circle), for CD4+ effector subset (n = 13 parameters). (D) Hierarchically clustered heatmap represent the difference in percent for each individual immune cell trait between HD-MS twin compared to SCNI-MS twin within each pair for the CD4+ effector subset. (E) Independent validation cohort consisting of healthy controls (HD, n = 71; turquoise), CIS (n = 55, light olive) and treatment naïve MS patients (MS, n = 60; olive) of selected CD4+ effector parameters. Statistical significance was evaluated by linear mixed models, as described in Materials and Methods and Mann–Whitney U test or unpaired t test; *P ≤ 0.05; **P ≤ 0.01; ns: not significant.
B cells (i.e., 18%) (24). Although the reasons for this differential influence have not been fully elucidated, the strong association between HLA background and T cell receptor repertoire may contribute to these findings (28). In addition to these observations highlighting the impact of genetics, a large twin study in healthy individuals revealed that also nonheritable influences have a distinct impact on immune traits as illustrated by the profound effect of a latent CMV infection affecting more than 50% of all immune cell parameters investigated (25). These findings prompted us to include CMV serostatus as fixed factor in our modeling, and indeed, CMV serostatus explained up to 17% of immune cell parameter variance in our dataset. Along this line, a Belgian survey using a large cohort of healthy individuals provided evidence of a profound impact of local environmental influences as reflected by cohabitation of nonrelated individuals (i.e., parents), which resulted in a 50% reduction in immunological variation between nonrelated individuals (18). Some of the nonheritable influences might lead to MS-associated epigenetic changes in immune cells, but these have yet to be characterized in detail (29).

In our study, the observed strong effect of twinship is most likely driven by a combined effect of genetic and shared environmental influences because of the fact that all twin pairs in our study were raised together until adulthood. Potentially relevant shared environmental influences range from infections to dietary influences and their respective effects on individual microbiome composition to lifestyle factors, such as exercise, smoking, and alcohol consumption. Importantly, many of these factors not only have been identified as modulators of immune cell traits but have also been implicated in the development of autoimmunity. Particularly in MS, there is evidence that at least some environmental influencing factors not only impact disease incidence and disease course, but also elicit changes in immune signatures (30, 31).

An important aspect that might have contributed to the moderate influence of MS at the level of the whole cohort is the relative heterogeneity of the MS-affected cotwins with regard to disease duration (range 0.5 to 45 y), disease course (relapsing form 67%, progressive form 33%), disease severity (expanded disability status scale [EDSS] range 0 to 9.5), as well as immune modulatory treatment (Table 1). In light of the obvious difficulties in the recruitment of monozygotic twins discordant for MS-Related Signatures in Cotwins with Prodromal MS. To overcome this limitation, we chose a complementary approach to investigate the impact of autoimmune immunology on immune signatures. Because the evolution of autoimmune diseases such as MS is a gradual process, we hypothesized that an analysis of MS-related effects on immune traits would be most promising at the earliest stages of the disease as this avoids any confounding effects related to disease stage and treatment. This concept of a gradual evolution has gained much interest in recent years (32, 33). In our cohort, the clinically healthy cotwins have a maximum familial risk for developing MS (34). Indeed, we were able to identify a subcohort of clinically healthy cotwins who showed signs of subclinical neuroinflammation (8). This offered us the unique possibility to study potential MS-related effects on each immune trait by comparing the degree of correlation in twin pairs who were clinically and subclinically discordant, to those where the healthy twin exhibited subclinical signs of prodromal MS. By this approach, we were indeed able to identify a pattern of potential MS-associated adaptive immune traits, especially within CD4+ effector T cell subsets. This finding supports the concept that MS is initiated and orchestrated by antigen-driven T cells (6, 7).

Notably, we had the opportunity to analyze flow cytometry data from a second independent cohort of untreated patients with either clinically isolated syndrome or early RRMS to evaluate whether the immune traits identified in our prodromal MS approach might be differentially expressed in early untreated MS patients. Indeed, several of our identified immune traits were found to be differentially expressed, in particular within the Th1 and Th17 effector populations. Despite using distinct approaches to study MS-related changes of immune signatures in incipient disease in both cohorts, our analysis revealed that very early disease stages were particularly associated with changes in peripheral immune signatures.

Interestingly, some of these immune traits identified in our systems approach had already been implicated in MS pathophysiology, either based on preclinical findings from animal models of MS or differential expression patterns using focused hypothesis-driven approaches in smaller cohorts of MS patients (35). For example, Th17 cells and a specific subset, termed Th17.1 cells, have both been shown to be highly encephalitogenic in EAE models and found to be increased in the peripheral blood of early MS patients (36–38). However, in contrast to these previous approaches, which investigated single individual candidates in a hypothesis-driven fashion, we now employed an unbiased approach that allowed us to identify MS-associated signatures in distinct immune cell populations of our twin cohort. Our findings are in line with a recent study employing a combination of single-cell mass cytometry (CyTOF) with machine-learning algorithms to identify a T-helper cell signature in MS patients (39).

It is interesting to note that our approach did not reveal a significant impact of MS on B cell traits. This finding seemingly contradicts the increasingly acknowledged role of B cells in MS highlighted by the success of B cell-depleting treatment strategies both in RRMS and primary progressive MS (40, 41). However, we acknowledge that our flow cytometry panels captured a higher number of B cell as compared to B cell traits, which in contrast to B cell traits include functionality assessments, resulting in a higher power for detection of alterations in the T cell compartment. Furthermore, our analysis in Fig. 2B revealed that B cell traits might be particularly susceptible to immune treatment effects. It should be pointed out that our regression analysis also suggested more pronounced MS effects on a few selected B cell traits, for example regulatory B cells and memory B cells. In addition, for selected antigen-presenting monocyte and DC populations, an individually higher MS influence could be noted, which is in line with other publications highlighting the potential role of specific myeloid cell subsets in CNS autoimmunity (42, 43).

The observed predominance of MS-associated adaptive immune traits in our correlation approach might at least partly be related to the fact that our phenotyping strategy overall covered more adaptive (102 traits) than innate (39 traits) immune traits. We therefore cannot rule out that a more detailed investigation of innate immune traits might reveal additional MS-associated features in these populations, since innate populations are increasingly acknowledged as key players in the pathogenesis of MS (44–46).

From a more general perspective, our approach illustrates that a stringent control for interindividual heterogeneity by use of a twin design combined with a focus on very early disease stages, thus minimizing disease-related heterogeneity, represents a powerful strategy to elucidate changes of the peripheral immunome in human autoimmunity in a comprehensive fashion. By analyzing subjects with subclinical (prodromal) signs of neuroinflammation we were able to uncover MS-associated immune signatures that would otherwise have escaped detection. We are aware of the relatively small sample size, but this is outbalanced by the distinct advantages of our twin approach. It should be mentioned that, albeit our deep-immune phenotyping panel encompasses a large number of immune traits, we cannot rule out that it does not fully capture more sophisticated functional immune
features, such as antigen recognition, functional exhaustion, active suppression, or other tolerance mechanisms.

In particular, in vitro stimulation of T cells with relevant target autoantigens and ex vivo staining with antigenic-peptide-MHC multimers might have revealed stronger MS-associated "signatures." However, apart from technical hurdles, validated target antigens are lacking in MS (6, 7). This contrasts with other autoimmune diseases of the CNS where the target antigens are well established, for example, neumyelitis optica spectrum disease (characterized by anti-aquaporin-4 antibodies) and myelin-oligodendrocyte glycoprotein (MOG)-antibody associated autoimmune disorder (characterized by anti MOG-antibodies) (47).

Another limitation is that with this approach we analyzed only blood and not CSF. While CSF analysis likely could provide important additional information, it poses practical and ethical obstacles because it would imply systemic invasive CSF sampling from healthy individuals. From a scientific point of view, however, analysis of immune signatures in secondary lymphatic structures as well as in CSF from our twin pairs would be highly interesting as these might more closely reflect target-organ and autoimmunity-associated changes of immune cell traits. Indeed, in a small cohort of monozygotic twins with prodromal MS, single-cell RNA sequencing of CSF cells revealed alterations in the adaptive immune compartment, such as clonal expansion of tissue-resident memory CD8+ T cells, plasma blasts, as well as CD4+ T cells, suggesting an involvement of all major components of the adaptive immune system (8).

Taking these data together, by controlling for the dominant influence of genetic and local environmental factors, and by focusing on incipient MS, we were able to reveal peripheral MS-related immune signatures pointing to a pivotal role of effector CD4+ T cells during disease initiation. Our study underscores the crucial importance of genetic and environmental heterogeneity as an obstacle in the search for disease-related immune signatures. Our in-depth analysis of healthy as well as cotwins with prodromal MS helped us to identify patterns of MS-associated immune alterations, some of which could be corroborated in an independent cohort of early, untreated RRMS patients. In future studies it will be of great interest to correlate such MS-linked variations of immune phenotypes with the emerging genetic and epigenetic risk profile of MS.

Materials and Methods

Study Cohorts. Twins were recruited between May 2012 and February 2016 in Germany. Inclusion criteria for study participation were met if only one cotwin of monozygotic twins had an MS diagnosis according to the revised McDonald criteria (10, 21). Exclusion criteria were infection, as well as treatment with antibiotics or high dose intravenous glucocorticosteroids within 3 mo prior to sampling. Monozygotic twin pairs clinically discordant for MS visited the outpatient department at the Institute of Clinical Neuroimmunology in Munich for a detailed interview, neurological examination, MRI investigations, blood sampling, and optional CSF sampling. To confirm MS diagnosis, medical records including MRI scans were obtained and reviewed. Exclusion criteria for the present analysis were a current disease modifying treatment (DMT) with fingolimod, alemtuzumab, dimethylfumarate, or teriflunomide, or other tolerance mechanisms. Cotwins were stained and measured on the same day to minimize acquisition variability. To label cell surface molecules, cryoconserved PBMCs were reconstituted in PBS with 5% BSA and 2 mM EDTA containing fluorochrome-conjugated antibodies and incubated for 30 min at 4 °C or 37 °C. For intracellular staining, cells were fixed and permeabilized for 20 min at 4 °C with Cytofix/CytopermTM working solution (biotiniscience) according to the manufacturer’s instructions, and subsequently stained with intracellular fluorochrome-conjugated antibodies for 30 min at 4 °C. To determine intracellular cytokine secretion, PBMCs were restimulated in X-vivo (Lonza) at 1 × 10^6 cells/mL in RPMI (Lonza) with Leukocyte Activation beads (Cytomix, Brefeldin A; BD Pharmingen) for 4 h at 37 °C with 5% CO2. Working antibody concentrations are depicted in Dataset S3. All flow cytometry data were acquired by Navios (twin cohort) and Cytosent (Beckman Coulter, validation cohort) flow cytometers and analyzed by Kaluza analysis software version 1.5 (Beckman Coulter) and GraphPad Prism 6. Data analysis was performed by two independent scientists (four-eyes principle).

Single-Cell Multiparameter Visualization. We used an unbiased approach to visualize flow cytometry data on a multiparameter single-cell basis. We analyzed 13 flow cytometry panels consisting of 9 to 10 fluorescence markers were measured (cellular functional immune phenotyping matrix). Gating strategies and identification of populations are depicted in Dataset S2. All samples were stained and measured over a short period of 4 wk by two experienced technicians on the same flow cytometer. Cotwins were stained and measured on the same day to minimize acquisition variability. To label cell surface molecules, cryoconserved PBMCs were reconstituted in PBS 0.5% BSA/2 mM EDTA containing fluorochrome-conjugated antibodies and incubated for 30 min at 4 °C or 37 °C. For intracellular staining, cells were fixed and permeabilized for 20 min at 4 °C with Cytofix/CytopermTM working solution (biotiniscience) according to the manufacturer’s instructions, and subsequently stained with intracellular fluorochrome-conjugated antibodies for 30 min at 4 °C. To determine intracellular cytokine secretion, PBMCs were restimulated in X-vivo (Lonza) at 1 × 10^6 cells/mL in RPMI (Lonza) with Leukocyte Activation beads (Cytomix, Brefeldin A; BD Pharmingen) for 4 h at 37 °C with 5% CO2. Working antibody concentrations are depicted in Dataset S3. All flow cytometry data were acquired by Navios (twin cohort) and Cytosent (Beckman Coulter, validation cohort) flow cytometers and analyzed by Kaluza analysis software version 1.5 (Beckman Coulter) and GraphPad Prism 6. Data analysis was performed by two independent scientists (four-eyes principle).

Determination of Biological Distance. Conventionally gated populations of the sub collectives CD4+ were divided into activated, effector, and other subsets (depicted in Dataset S4) according to their immunological phenotype (total
number of investigated populations: CD4+ activated subset, n = 6; CD4+ other subset, n = 7; CD4+ effector subset, n = 13). Frequency values per population were normalized and clustered by the t-SNE algorithm using the Rtsne package in R. Biological distance was determined based on the Euclidean distance of t-SNE coordinates (16, 18, 58). To account for clustering variability and assess analysis robustness, the analysis was performed three times (without a seed) and the mean was used to determine the biological distance. Only twin pairings with complete datasets per parameter were included into this analysis. Statistical significance was determined with GraphPad Prism 6 by the Mann–Whitney U test: *p < 0.05; **p < 0.01.

Regression Analysis. Regression analyses were performed using SAV v9.4 (SAS Institute) and R v3.6.1 (59). For each immune parameter, a linear model was fitted including age, gender, CMV, and disease status (MS yes/no) as fixed effects as well as a random intercept in order to account for the correlation within twin pairs (SAS, proc mixed). Based on the mixed models, marginal and conditional R²s were computed according to Nakagawa and Schielzeth (60). The authors indicate the proportion of variance explained based on the fixed effects (marginal R²) and the fixed plus the random effects (conditional R²) (i.e., also accounting for the correlation within pairs of twins). ICCs were calculated by dividing the variance estimate of the random effect by the sum of this variance and the residual variance. In addition, correlation within twin pairs for each parameter. Additionally, the proportion of variance explained by the different fixed effects was estimated using the relaimpo package (lmig metric) (61). In order to account for correlation within twin pairs, a workaround for clustered data by specifying a survey design was used as presented in the package manual.

Mixed-model analyses were repeated for the two separate twin cohorts with healthy cotwins and SCNI cotwins. The ICCs from these models (adjusted for age, gender, and CMV) were compared using a mixed model accounting for the dependence of immune parameters. Therefore, the correlation of immune parameters was performed based on the whole cohort using the marginal residuals from linear mixed models (i.e., accounting for age, gender, and CMV). Because of the correlation of residuals within twin pairs, the residuals were averaged for each pair of twins (62). The correlation between immune parameters was then estimated using Spearman’s rank correlation coefficient. The sample correlation matrix was implemented in the mixed model by applying the lm4qtl package in order to specify the covariance matrix for the random effects (63). No adjustment for multiple testing was applied (i.e., all significance levels are to be understood as local significance levels). Therefore, inference procedures are intended to be exploratory (hypotheses generating), not confirmatory.

Data Availability. All study data are included in the article, the SI Appendix, and Datasets S1–S8. Full protected access of flow cytometry data will be granted to collaborating scientists who are willing to complete an “MS-TWIN Data Access Agreement” and an “MS-TWIN Data Access Application form.”

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