CD4+Foxp3+ T Regulatory Cells Promote Transplantation Tolerance by Modulating Effector CD4+ T Cells in a Neuropilin-1-Dependent Manner

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Several mechanisms of immune suppression have been attributed to Foxp3+ T regulatory cells (Treg) including modulation of target cells via inhibition of cell proliferation, alteration of cytokine secretion, and modification of cell phenotype, among others. Neuropilin-1 (Nrp1), a co-receptor protein highly expressed on Treg cells has been involved in tolerance-mediated responses, driving tumor growth and transplant acceptance. Here, we extend our previous findings showing that, despite expressing Foxp3, Nrp1KO Treg cells have deficient suppressive function in vitro in a contact-independent manner. In vivo, the presence of Nrp1 on Treg cells is required for driving long-term transplant tolerance. Interestingly, Nrp1 expression on Treg cells was also necessary for conventional CD4+ T cells (convT) to become Nrp1+Eos+ T cells in vivo. Furthermore, adoptive transfer experiments showed that the disruption of Nrp1 expression on Treg cells not only reduced IL-10 production on Treg cells, but also increased the frequency of IFNγ+ Treg cells. Similarly, the presence of Nrp1KO Treg cells facilitated the occurrence of IFNγ+CD4+ T cells. Interestingly, we proved that Nrp1KO Treg cells are also defective in IL-10 production, which correlates with deficient Nrp1 upregulation by convT cells. Altogether, these findings demonstrate the direct role of Nrp1 on Treg cells during the induction of transplantation tolerance, impacting indirectly the phenotype and function of conventional CD4+ T cells.

Keywords: Neuropilin-1, tolerance, Treg cells, immune regulation, transplantation

INTRODUCTION

Foxp3+ T regulatory (Treg) cells are an important population of leukocytes that control immunity, mainly by dampening effector T cell responses. Many studies have described the mechanisms by which Treg cells carry out their function, such as IL-2 deprivation, secretion of cytotoxic granules (granzyme/perforin), metabolic disruption, secretion of anti-inflammatory cytokines, and release of extracellular vesicles (exosomes) (1, 2).

In addition to their capacity to suppress immune responses, Treg cells had become an interesting target for cell therapy, due to the increasing number of diseases associated with malfunction and
over-reactivity of the immune system, such as autoimmunity and transplant rejection (3, 4). The current paradigm is based on the premise that immune tolerance to allogeneic transplant is broken by an imbalance of Treg cells over T effector cells. The infusion of Treg cells considerably increases graft survival in transplanted animals (4, 5), and clinical trials of the administration of Treg cells into patients have demonstrated safety but variable efficacy (6). Understanding Treg cell biology and its mechanisms of immune suppression may improve the potential and use of Treg cells as therapeutic agents.

A few years ago, Neuropilin-1 (Nrp1) was described as a potential Treg cell marker. Nrp1 is a transmembrane co-receptor with affinity for a variety of ligands, all involved in physiological processes, such as angiogenesis, axonal guidance, or immune synapses (7). In the immune system, Nrp1 is expressed mainly by dendritic cells, Natural Killer (NK) and Treg cells (8–11). Initially, the function described for Nrp1 was to stabilize the interaction between cells during antigen presentation through homotypic interactions (12, 13). However, other studies suggested later that Nrp1 contributes to the function, phenotypic stability, and survival of Treg cells in tumors (14).

Several reports correlate Nrp1 expression on T cells with a state of immune tolerance (14–19), which has been demonstrated in the transplantation context both in patient biopsies and experimental models (20–22). In addition, Nrp1-deficient or Nrp1KO Treg cells are not capable of exerting suppressive function through a semi-porous membrane; and the same phenomenon was observed when using wild type Treg cells in the presence of anti-Nrp1 blocking antibodies (14).

We previously described that conventional CD4+ T cells (defined as CD4+CD25-Nrp1-Foxp3-cells or convT) up-regulate Nrp1 expression during allograft rejection. Interestingly, in the tolerogenic condition in which Nrp1+Foxp3+ Treg cells are co-transferred with convT cells, a larger frequency of Nrp1+Eos+ convT cells was observed suggesting that Nrp1+Treg cells could modulate the phenotypic signature of convT cells (22), leading to the generation of T cells with modulatory effects.

Based on these antecedents, we hypothesized that convT cells gain Nrp1 and Eos in an Nrp1+Treg cell-dependent manner to favor immune suppression. Using Nrp1 conditional knocked out mice; we demonstrate that Nrp1KO Treg cells are deficient in exerting suppressive activity in a contact-independent manner. Even more, when Treg cells lack Nrp1, convT cells are unable to up-regulate Nrp1 and Eos expression favoring the appearance of type-1 T helper (Th1) cells. Accordingly, the frequency of IL-10+Treg cells is negatively affected, which correlates with the inability to induce long-term tolerance. Lastly, we demonstrate that Treg cells-modulated convT cells also gain the ability to suppress ex vivo T cell proliferation, which is affected if co-transferred Treg cells lack Nrp1. Hence, we demonstrate that Treg cells drive immune tolerance by modulating the phenotype and function of convT cells in an Nrp1-dependent manner.

RESULTS

The Lack of Nrp1 on Treg Cells Is Not Involved in Treg-Phenotypic Signature

In 2015, our group reported that convT cells transferred into skin-transplanted animals gain Nrp1 expression (from 0 to ~35%). This induction was highly increased when convT cells were co-transferred with Nrp1+Treg cells (22). To clarify whether this process depends on Nrp1 expressed specifically on Treg cells, we obtained Foxp3Cre/YFP and Nrp1flx/flox mice to generate Nrp1-deficient or Nrp1KO Treg cells, which are conveniently detectable by flow cytometry based on the expression of YFP (23, 24). First, we tested the phenotype of T cells from different organs/tissue of Foxp3Cre/YFP (wild type, wt control), Foxp3Cre/YFP Nrp1flx/+ (het) and Foxp3Cre/YFP Nrp1flx/flx (Nrp1KO Treg) offspring. As expected, we found that deletion of Nrp1 only occurs on Treg cells, as seen in peripheral lymph nodes (pLN), spleen (Spl), and blood cells (~75% on wt and het, and <1% on Nrp1KO) (Figure 1A). In the case of convT cells, we find a partial decrease in Nrp1 expression only in the Spl, which could correspond to antigen experienced ex-Treg cells (Foxp3-T cells), since convT cells were considered and gated as Foxp3-cells.

In addition, CD8+ T cells express very low levels of Nrp1 (<5%), which did not change either in het or Nrp1KO Treg cells (Supplemental Figure 1A) (15). We also analyzed the expression of Treg cell-associated markers such as Eos (25, 26), CD49b (27), CD25 (28), Helios (23), CD103 (29–31), and CD73 (32, 33) along with T-bet, GATA-3, and RORγT as lineage transcription factors for T helper (Th)-1, Th2, and Th17 cell subsets, respectively (34–36) on convT and Treg cells observing no differences among all three genotypes (Figures 1A,B). The absence of Nrp1 on splenic Treg cells obtained from Nrp1KO-animals was corroborated using western blot assay (Figure 1C).

Even more, we performed high-dimensional single cell data analysis by visualization of t-Distributed Stochastic Neighbor Embedding (t-SNE) algorithm (visSNE) (37, 38) on pLN, Spl and blood cells populations from all three mice genotypes (Figures 2A,B), searching for main cell subsets based on the expression of CD8, CD4, Foxp3, and CD19 (Supplemental Figure 1B). visSNE heat maps confirmed that Nrp1 depletion only occurs in the Foxp3+Treg cell compartment of Nrp1KO animals, whereas expression of Nrp1, Eos, CD49b, and CD25 remain mostly unchanged among the identified populations (CD19+ B cells, CD4+Foxp3-convT, and CD8+ T cells) (Figures 2B,C). Additionally, by performing spanning-tree progression analysis of density-normalized events (SPADE), we identified 11 spatially distinct immune cell populations with unaltered frequencies among wt, het, and Nrp1KO splenic cells (Figures 2D,E). Furthermore, the relative Mean Fluorescence Intensity (MFI) of each marker was calculated for each SPADE-on-visSNE cell population studied, highlighting the lack of
FIGURE 1 | Foxp3-targeted Nrp1 deletion only occurs on Treg cells and does not affect conventional T cell phenotype. Peripheral lymph nodes (pLN), spleen (Spl), and blood were harvested from Foxp3YFP/CreNrp1+/- (wt, dark green bars), Foxp3YFP/CreNrp1flox/flox (het, light green bars), and Foxp3YFP/CreNrp1flox/flox (Nrp1KO, white bars) animals. Leukocytes were obtained and tested for the indicated markers by flow cytometry. (A) Graphs show the frequency of Nrp1, Eos, CD49b, and CD25 on Foxp3+ Treg cells and conventional CD4+ T (convT) cells from wt, het, and Nrp1KO Treg cell-restricted mice. (B) Representative histograms showing the expression of indicated markers on Foxp3+ Treg cells from wt (dark green line), het (light green line), Nrp1KO (black line) mice; and FMO control (gray line). (C) Immunoblot of Nrp1 protein expression on wt and Nrp1KO Treg cells. For A, n = 3–4 animals per genotype, each circle represents one mouse, bars represent mean. Two-way ANOVA, * < 0.05; ** < 0.01; *** < 0.001; ns, not significant. Representative data of three independent experiments.
**FIGURE 2** Disruption of Nrp1 expression on Treg cells does not interfere with the acquisition of modulatory molecules. Organs and tissues were collected and processed as in Figure 1. viSNE analysis was run on 10,000 live single cells per sample using cell subtype markers. (A) Representative viSNE map of concatenated files of all samples, and concatenated individual populations from pLN, Spl, and blood samples. (B) viSNE map showing color-coded cell populations based on lineage marker expression gating. (C) Fluorescence intensity for specific markers from wt, het, and Nrp1KO spleen cells on viSNE map. (D) SPADE-on-viSNE map defining 11 spatially distinct splenic cell populations. (E) Relative proportions of immune cell populations defined in (D) from splenocytes. (F) MFI heat map of surface and intracellular markers expression on each immune cell subpopulation from splenic Treg-restricted wt (w), het (h), and Nrp1KO (k) mice. Representative data of at least three independent experiments.
Nrp1 expression on Nrp1KO Treg cells among different tissues (Figure 2F).

**Treg Cells Require Nrp1 Expression to Modulate Conventional CD4+ T Cell Phenotypic Signature**

After confirming typical Treg cell phenotype in all genotypes, we performed a suppression assay to test the modulatory function of Treg cells and acquisition of Nrp1 by convT cells in vitro. For this experiment we used congenically marked antigen presenting cells (APC, CD45.2+) Treg cells (CD45.2+) and CTV-stained convT cells (CD45.1+) to facilitate the tracking of convT cells under proliferation (Supplemental Figure 2A). As shown in Figure 3A, wt Treg cells suppress the proliferation of convT cells at 1:1, 1:2, and 1:4 Treg:convT ratios (~85, ~65, and ~45% of suppression, respectively); but both het and Nrp1KO Treg cells show slightly reduced suppressive activity, although these variations are not statistically different. These results suggest that Nrp1 is not fully required for Treg cells acting in a contact-dependent manner (Figure 3B). On the other hand, we checked the phenotype of convT cells by looking at Nrp1 expression. As depicted in Figures 3C,D, convT cells acquired Nrp1 expression in a dose and Treg-dependent manner as Nrp1KO Treg cells were able to allow convT cells to become Nrp1+ to a lesser extent (for instance, at 1:1 ratio we observe ~5% of Nrp1+ conv T cells without Treg cells, ~20% of Nrp1+ convT cells with wt Treg cells, ~15% of Nrp1+ convT cells with het Treg cells and ~12% of Nrp1+ convT cells with Nrp1KO Treg cells). This data supports our previous observation, in which convT cells up-regulated Nrp1 during allograft rejection but when convT cells were co-transferred with Treg cells, the expression of Nrp1 was enhanced in addition to the induction of transplant tolerance (22). At the same time, Nrp1-expressing Treg cells (wt and het) decrease Nrp1 expression in a dose-dependent manner when co-cultured with convT cells at the different ratios, Figure 3E. Conversely, convT cells co-cultured with wt Treg cells did not gain Eos expression in vitro, which could be a result of the experimental timing (3 days for in vitro assay vs. 20 days for in vivo assay) (Supplemental Figures 2B,C).

Since we did not observe a robust difference between the suppressive activity of wt, het, and Nrp1KO Treg cells, we repeated the suppression assay but using a contact-independent setting (transwell system) (Supplemental Figure 3). In this case, we observed differences in the suppressive function of wt and Nrp1KO Treg cells, where Nrp1KO Treg show reduced suppressive activity, indicating that Nrp1+ Treg cells produce factors with modulatory function, which are altered in Nrp1KO Treg cells, Figures 3F,G. Even more, convT cells also up-regulated Nrp1 expression when wt Treg cells, and not Nrp1KO Treg, were placed at the top chamber (~17% vs. ~9% of Nrp1+ convT cells, respectively, Figure 3H). Thus, the presence of Nrp1 on Treg cells modulates the phenotype of convT cells and contributes to T cell suppression in vitro.

**Nrp1KO Treg Cells Cannot Induce Long-Term Transplant Tolerance and Fail to Modulate Conventional CD4+ T Cells in vivo**

Next, we tested the role of Nrp1 on Treg cell function using an in vivo skin transplantation model, in which C57BL/6 × Balb/c (F1) skin is grafted onto RAG-KO mice previously administered with convT cells (Supplemental Figure 4A). Using this approach, we observed complete transplant rejection by day 20 post-surgery, but skin graft tolerance when convT cells are co-transferred with Nrp1+ Treg cells (22). In the current study, we carried out long-term skin transplant experiments using wt, het, and Nrp1KO Treg cells and found that wt Treg cells induce ~60% of transplant survival and only ~20% for the groups receiving het and Nrp1KO Treg cells (Figure 4).

Furthermore, we performed 20 days-long experiments, for which skin graft-draining lymph nodes (dLN) were removed and the number and phenotype of T cells was studied. All three Treg cell genotypes were able to reduce cellularity in the dLN of allografted animals (Supplemental Figure 4B). Additionally, we analyzed dLN cell phenotype by flow cytometry, discriminating clearly between convT cells (CD45.1+) and Treg cells (CD45.2+) (Supplemental Figures 4C,D). As depicted in Figures 5A,C, convT cells gained Nrp1 and Eos expression, which seems to occur in a Treg cell-dependent manner because ~30% of convT cells became Nrp1+ when co-transferred with wt Treg cells in comparison with ~15% or ~12% when co-transferred with het or with Nrp1KO Treg cells, respectively, Figure 5B. Moreover, ~20% of convT cells gained Eos expression when co-transferred with wt Treg cells in contrast to ~10% for the case of het or Nrp1KO Treg cells, Figure 5D. Importantly, Eos expression remain unaltered regardless of Treg cells genotype, Supplemental Figures 4D–G. These observations confirmed our previous study in which we described that CD4+ T effector phenotype is modulated by Nrp1-expressing Treg cells.

To further investigate Nrp1KO Treg cells, we studied Treg cell cytokine production finding that ~20% of either het or Nrp1KO Treg cells were IFNγ+ in contrast to~10% of wt IFNγ+ Treg cells, Figures 6A,B. No differences were observed among the frequencies of wt, het, or Nrp1KO Treg cells either IL-17A+ or IFNγ+IL-17A+. When we tested for the production of the anti-inflammatory cytokine IL-10, we found that ~5% of het and Nrp1KO Treg cells were IL-10+ compared with ~8.5% of wt Treg cells, Figures 6C,D. Therefore, Nrp1-deficiency on Treg cells during allograft rejection negatively affects IL-10 production but favors IFNγ production, which may explain the reduced % of transplant survival of grafted animals treated with Nrp1KO Treg cells. To corroborate this, we analyzed IL-10 levels on convT-and-Treg in vitro culture supernatants, observing decreased IL-10 production when convT were co-cultured with Nrp1KO Treg cells, compared with wt
FIGURE 3 | Nrp1 is required for immune-cell suppression and modulation of conventional CD4+ T phenotype. Responder convT cells were sorted, CTV-labeled and cultured with Mitomycin-C treated-APCs plus soluble anti-CD3 antibody, alone or in the presence of sorted Treg cells. (A) Histograms showing proliferation of convT cells activated alone (gray) or at different Treg:convT ratios (1:1, 1:2, and 1:4) of wt (dark green), het (light green), or Nrp1KO (black) Treg cells. (B) Percentage of suppression activity was calculated as described in Materials and Methods. Representative data of at least three independent experiments. Squares represent mean ± SEM. (C) Representative dot plots showing Nrp1 expression on convT cells activated as in (A). (D) Pooled data depicting Nrp1 expression on CD45.1+ convT cells (Continued)
FIGURE 3 | cultured with Treg cells from the three indicated genotypes. Bars represent mean ± SEM. Data is representative of at least two independent experiments.

(E) Pooled data depicting Nrp1 expression on CD45.2+ Treg cells before and after co-culture with convT cells at aforementioned different Treg:convT ratios. PS: post-sorting expression of Nrp1 on Treg cells. (F) Histograms showing proliferation of convT cells activated alone (gray) of at different Treg:convT ratios (1:2 and 1:4) of wt (dark green), het (light green), or Nrp1KO (black) Treg cells from transwell experiments (described in detail in Materials and Methods section and Supplemental Figure 3). (G) Percentage of suppression activity was calculated as described in Materials and Methods. Representative data of two independent experiments performed in triplicates. (H) Pooled data depicting Nrp1 expression on CD45.1+ convT cells cultured with wt or Nrp1KO Treg from transwell experiments. Bars represent mean ± SEM. Data is representative of two independent experiments performed in triplicates. For (B,D–E,H), Paired T-test; * < 0.05; ** < 0.01 were used; ns, not significant. For (G) * < 0.05 for wt vs. Nrp1KO (both 1:2 and 1:4 ratios).

FIGURE 4 | Nrp1KO Treg cells are unable to exert long-term skin transplantation tolerance. Conventional CD4+ T cells were sorted and adoptively transferred alone or with sorted Foxp3/YFP+CD45.2+ Treg cells into RAG-KO recipient mice. Next day, mice were transplanted with F1 semi-allogeneic skin grafts (day 0) and graft survival was monitored over time. Long-term skin allograft survival was compared between mice receiving wt (dark green), het (light green), or Nrp1KO (black) Treg cells. Lines represent graft survival (%), log-Rank test: *p < 0.05 between “convT cells” and “wt”; *p < 0.05 between “convT cells” and “het”; *p > 0.05 between “convT cells” and “Nrp1KO”; *p > 0.05 between “wt” and “het”; *p < 0.05 between “wt” and “Nrp1KO”.

FIGURE 5 | Nrp1+ Treg cells are required for the acquisition of Nrp1 and Eos expression by conventional CD4+ T cells during skin transplantation acceptance. Mice were transplanted as described in Figure 4. At day 20 post-surgery, graft-draining lymph nodes were harvested and cells were analyzed for Nrp1 and Eos expression by flow cytometry. Representative contour plots depicting (A,B) Nrp1 and (C,D) Eos expression on gated live CD4+CD45.1+ convT cells. (B–D), bars represent mean and each circle represents one mouse. Unpaired T-test, * < 0.05; ** < 0.01; ns, not significant; PS, post sorting. Representative data of at least two independent experiments.

Treg (Figure 6E). According to our previous results, convT cells upregulate Nrp1 expression when cultured with wt Treg but not with Nrp1KO Treg cells (~30% Nrp1+ convT vs. ~20% Nrp1+ convT cells cultured with Nrp1KO Treg or alone, Figure 6F). Notably, the addition of exogenous IL-10 to the culture appeared to rescue the gaining of Nrp1 expression by convT cells supporting the notion that decreased IL-10 production contributes to the impaired suppressive
function of Nrp1KO Treg and their immune modulation on convT cells.

When we analyze the convT cell subset, we found ~40% of IFNγ+CD4+ convT cells in the rejecting group; this frequency was significantly reduced to ~15% in the presence of wt Treg cells, but only to ~25% of convT cells in the presence of het or Nrp1KO Treg cells, Figures 7A,B. The frequencies of IL-17+CD4+ T and IFNγ+IL-17+ CD4+ T cells were not changed in the presence of Nrp1KO Treg cells, Figures 7C,D. Strengthening this observation, and the fact that wt Treg cells modulate convT cells phenotype, we sorted out convT cells from transplanted animals that received either wt or Nrp1KO Treg cells, and tested them for in vitro inhibitory function (Figure 7E), finding that the former also renders suppressive function, as
shown in the ex vivo suppression assay (~55% proliferation of freshly isolated CD45.2+ convT cells co-cultured with wt Treg-modulated CD45.1+ convT cells vs. freshly isolated convT cells co-cultured with effector T cells from rejecting animals), Figures 7E,G. Accordingly, convT cells become less suppressive when Treg cells are defective in Nrp1 expression (~105% proliferation of fresh CD45.2+ convT cells co-cultured with “Nrp1KO Treg-modulated convT cells”).

Altogether, our findings indicate that Treg cells require Nrp1 to modulate the function of convT cell and to exert optimal suppressive activity in vivo.

DISCUSSION

Regulatory T cell malfunction has been widely associated with increased inflammatory immune responses. Understanding Treg cells biological processes and mechanisms of suppression are pivotal for recognizing new targets for therapy. Nrp1 has been previously proposed as a cell marker for thymic-derived Treg cells (39); but its expression has also been described on T cells during allogeneic skin graft rejection (22), sepsis (40), IL-10 deficiency (41), and anti-tumor immune responses (42, 43). Moreover, Nrp1 deficiency on Foxp3+ Treg cells has been associated with lack of immune suppression, predominantly affecting tumor growth (14, 15, 44) and worsening EAE severity (45). In this work, we focused on analyzing the role of Nrp1 specifically on Treg cells during the induction of skin transplant tolerance. In one of our previous studies we demonstrated that Nrp1+ Treg cells drive transplantation tolerance, proposing the modulation of conventional CD4+ T cell phenotype as a possible mechanism (22). To get more insight regarding this observation, we obtained conditional knockout animals in which the lack of Nrp1 expression is restricted to Foxp3+ Treg cells (24). First, we extensively analyzed the phenotype of T cells in these genetically modified mice, including Foxp3+ Treg cells, finding no aberrant expression of Treg cell-associated markers in animals containing Nrp1KO Treg cells compared to controls (Figures 1, 2). Next, we designed in vitro experiments to evaluate the function of Nrp1KO Foxp3+ Treg cells and their capacity to modulate convT cell phenotype. The regulatory ability of Nrp1KO Foxp3+ Treg cells was evaluated in contact-dependent and—independent assays, observing that Foxp3+ Treg cells require Nrp1 to inhibit T cell proliferation in a contact-independent manner, suggesting that Nrp1+ Treg cells secrete modulators to exert their suppressive function (Figure 3). Because we obtained a defect in cell contact-independent suppression when using Nrp1KO Treg cells, and Delgoffe et al. reported that the inclusion of anti-Nrp1 blocking antibody in a transwell suppression assay negatively affected the suppressive capacity of Nrp1+ Treg cells (14), it is reasonable to propose that Nrp1+Foxp3+ Treg cells may secrete Nrp1+ extracellular vesicles (EV) to “deliver” Treg cells’ modulatory effects to convT cells. In this regard, several authors have published that Treg cells are able to produce EV (containing modulatory molecules) as a novel manner to inhibit CD4+ T cell effector function (46–48). Furthermore, we also observed in the in vitro experiments that convT cells gained Nrp1 in a dose dependent manner according to the ratios used with Nrp1+ Treg cells (Figures 3C,D). In other words, we obtained higher % of Nrp1+ convT cells when co-cultured with higher number of Nrp1+ Treg cells, resulting in lower frequencies of Nrp1+ convT cells when Nrp1KO Treg cells were added. Even more, Nrp1+ Treg cells decreased Nrp1 expression while convT cells gained it (Figures 3D,E). This phenomenon was first reported by our group using an in vivo model of transplantation and in this current report we corroborated the observation in vitro in addition to proving that it occurs in Nrp1+Foxp3+ Treg cell-dependent manner (Figure 3). Hence, it is conceivable to propose that Nrp1+ Treg cells could secrete Nrp1+ EV to target convT cells; in this regard, data from our laboratory indicates that wt Treg cells secrete EV containing Nrp1 in their membrane and that these EV can modulate convT cells phenotype and function, in comparison to EV obtained from Nrp1KO Treg cells (manuscript in preparation).

Using a murine model for allograft transplantation, we demonstrated that Nrp1 is required by Foxp3+ Treg cells to facilitate long-term tolerance (Figure 4). Interestingly, when we studied the phenotype of wt and Nrp1KO Treg cells in this in vivo setting, we did not find changes in Foxp3 or Eos expression (Supplemental Figures 4D–F). The transcription factor Eos functions as a co-repressor of Foxp3, preventing the expression of convT cell-related genes in Treg cells (26); and it has been described that “Eos-labile” Treg cells display a more inflammatory/helper phenotype (25). On the contrary, Rieder et al. have reported that convT cells from Eos−/− animals produce less IL-2 and show a malfunctioning CD25/STAT5 signaling pathway upon in vitro activation, but under in vivo inflammatory conditions Eos−/−T cells become high producer of IL-17 (49). In this work, we found that Nrp1 deficiency did not modify Foxp3 or Eos expression on Foxp3+ Treg cells under homeostatic conditions and in our model of skin graft transplantation, supporting that graft rejection was not due to instability in the expression of Foxp3 or Eos. Complementing Treg cell phenotypic analysis, we found that Nrp1KO Treg cells up-regulate IFNγ (Figure 6), supporting that the Nrp1 signaling pathway is required by Treg cells to secrete the appropriate cytokines (44). Gao et al. showed that Nrp1low CD25+CD4+ Treg (Treg expressing low levels of Nrp1) purified from septic mice secreted lower amounts of IL-10 (17), and another study using tumor-harboring II10KO (IL-10 deficient)-mice showed decreased Nrp1 expression in tumor-infiltrating Treg, impaired Nrp1+ Treg tumor accumulation and tumor protection function (41). While RNA sequence observations by Delgoffe et al. previously showed decreased II10 expression on Nrp1KO Treg (14), we have demonstrated here, for the first time, an impairment in IL10 production [both in vitro (in co-culture suppression assay) and in vivo (during allograft rejection), Figure 6] on Treg cells lacking Nrp1 expression. The gaining of Nrp1 expression by convT cells failed in the presence of Nrp1KO Treg (Figure 3) but, intriguingly, it could be reversed by adding exogenous IL-10 to the culture media, suggesting that Treg-mediated upregulation of Nrp1 on convT cells (and possibly other immunomodulatory proteins) could be dependent on IL-10 signaling.
FIGURE 7 | Nrp1+ Treg cells modulate conventional T cell phenotype and function. Mice were transplanted as described in Figure 4. At day 20 post-surgery, graft draining lymph node cells were obtained and stimulated with PMA and Ionomycin in the presence of Brefeldin A. Next, intracellular cytokine staining was performed and the expression of IFNγ and IL-17 on convT cells was determined by flow cytometry. (A) Representative dot plots showing the expression of IFNγ and IL-17 on CD4+CD45.1+ convT cells when transferred alone or with Treg cells from wt, het, or Nrp1KO Treg cells. Bar graphs depicting the frequencies of (B) IFNγ+CD4+ T, (C) IL-17+CD4+ T, and (D) IFNγ+IL-17+CD4+ convT cells when transferred alone (black) or with wt (dark green), het (light green), or Nrp1KO (white) Treg cells. Pooled data from two independent experiments, each circle corresponds to one mouse. Unpaired T-test, *< 0.05; **< 0.01; ns, not significant.

(E) Mice were transplanted as described in Figure 4. At day 15 post-surgery, graft draining lymph node cells were obtained, stained and live CD45.1+CD4+ convT cells were FACs-sorted and tested for suppression function over freshly isolated, CTV-stained CD45.2+CD4+ convT cells activated with MHC-II+APCs and soluble anti-CD3 for 72 h. (F) Histograms showing proliferation of fresh CD45.2+CD4+ convT cells activated alone (dark gray) or in the presence of control Foxp3+ wt Treg (orange) or CD45.1+CD4+ convT cells of dLN of RAG-KO allografted mice from Group (i) (convT alone, blue), Group (ii) (convT plus wt Treg cells, green), or Group (iii) (convT plus Nrp1KO Treg cells, light gray). (G) Percentage of proliferation of fresh CD45.2+CD4+ convT cells was calculated as described in Materials and Methods. Representative of at least three independent experiments. Bars represent mean ± SEM. Paired T-test, *< 0.05 was used.
MATERIALS AND METHODS

Mice
Six to 8 week old male and female mice were used. Foxp3<sup>Cre</sup>−<YFP> (24), Nrp1<sup>flox/flox</sup> (50), and RAG-KO mice were purchased from The Jackson Laboratory (Maine, USA), BALB/c (H-2<sup>b</sup>), C57Bl/6 (H-2<sup>b</sup>), F1 (C57Bl/6 × BALB/c, H-2<sup>a/b</sup>), and Foxp3<sup>3GFP</sup> (CD45.1 ± or Ly5.2) (51) mice were maintained under pathogen-free conditions at the animal facility located at Facultad de Medicina, Universidad de los Andes. All procedures were reviewed, approved and carried out according to the bioethics committee guidelines from Universidad de los Andes, and National Commission of Science and Technology (CONICYT).

Flow Cytometry
After cell viability staining with ZombieDye NIR (Biorad, CA, USA), cell suspensions were stained in 1X PBS with 5% heat-inactivated fetal bovine serum (Gibco, USA) with the following antibodies anti-: CD3e (clone 145-2C11), CD4 (clone RM4-5), CD8 (clone 53-63.7), CD19 (clone 6D5), CD25 (clone PC61), CD45.1 (clone A20), CD45.2 (clone 104), I-A/I-E (clone M5/114.15.2), CD103 (clone 2E7), CD49b (clone Hma2), CD73 (clone TV/11.8), Nrp1 (clone 3E12), all from Biolegend, CA, USA. Surface staining was performed for 30 min at 4°C. Intracellular staining was performed using anti-: Helios (clone ESB7C2, ThermoFisher Scientific), Foxp3 (clone FJK-16s), ROR<sub>yt</sub> (clone B2D), and the Fixation/Permeabilization Staining Kit (all from eBioscience, CA, USA), following manufacturer’s instructions. For cytokine expression analysis, cells were activated with 50 ng/mL PMA (Sigma) and 1 μg/mL Ionomycin (Sigma) in RPMI containing 10% FBS and Brefeldin-A (eBioscience, CA, USA) for 4 h. Cells were stained for surface markers, fixed in Fixation Buffer and washed with the Intracellular Staining Permeabilization Wash Buffer (all from Biolegend, CA, USA). Intracellular cytokine staining was performed using anti-: IFN<sub>γ</sub> (clone XMG1.2), IL-17A (clone TC11-18H10.1), and IL-10 (clone JES5-16E3). Cells were sorted on a FACSaria II (BD Biosciences, USA) and/or analyzed on a FACSCanto II (BD Biosciences, USA). Data analysis was performed using FlowJo software (CA, USA). To perform visualization of complex flow cytometry data, we used the Cytobank computational tool viSNE (visualization of t-Stochastic Neighbor Embedding) that generates a two-dimensional map in which cell distance represents the distance between cell parameters in high-dimensional space (37). Thus, cells that were phenotypically similar for the analyzed markers will be closer in a viSNE map (38, 52). To generate viSNE maps, samples were uploaded to Cytobank, live single cells were gated based on cell size and length and negative to Zombie Dye viability staining. Then, between 150,000 and 160,000 cell were subsampled from the data. After subsampling, viSNE was run at default parameters (1,000 iterations, random seed, perplexity = 30, theta = 0.5). viSNE maps were visualized using Cytobank interface, which was used to generate figures (color coding by marker expression levels).

Skin Transplantation
Tail skin (~1 cm<sup>2</sup>) from C57Bl/6 (syngeneic) or F1 (allogeneic) donors was transplanted onto the dorsal area of RAG-KO recipient mice. Survival of skin allografts was evaluated twice per week and grafts were considered rejected when 80% of the initial graft had disappeared or become necrotic.

Adoptive Transfer Experiments
Spleen and lymph node CD4+ T cells were pre-enriched using the CD4+ T Cell Isolation Kit, mouse (MiltenyiBiotec, CA, USA). CD4<sup>+</sup> Foxp3<sup>Cre</sup>− Nrp1<sup>−</sup> cells were sort-purified as CD4<sup>+</sup> T conventional cells from Foxp3<sup>3GFP</sup>CD45.1<sup>+</sup>reporter mice, and CD4<sup>+</sup> Foxp3<sup>Cre</sup>−/YFP<sup>+</sup> were FACS-sorted as Treg cells from Foxp3<sup>Cre</sup>/YFP<sup>CD45.2</sup> (wt), Foxp3<sup>Cre</sup>−/Nrp1<sup>flox</sup>/Foxp3<sup>Cre</sup>−/Nrp1<sup>flox</sup> (het), or Foxp3<sup>Cre</sup>−/Nrp1<sup>flox</sup>/Foxp3<sup>Cre</sup>−/Nrp1<sup>flox</sup>/Foxp3<sup>Cre</sup>−/Nrp1<sup>flox</sup> (Nrp1KO) mice. CD4<sup>+</sup> conventional T cells (1.5 × 10<sup>5</sup>) were intravenously (i.v.) injected alone or co-transferred with Treg cells (5 × 10<sup>4</sup>) into RAG-KO mice, 1 day before transplant surgery. Unless otherwise stated, 20 days post-transplant mice were euthanized and graft draining lymph nodes (dLNs) were extracted for flow cytometry analysis.

In vitro Suppression Assays
For contact-dependent assays, splenic CD4+CD25−Nrp1−Foxp3<sup>Cre</sup>− cells were sorted from Foxp3<sup>3GFP</sup>CD45.1<sup>+</sup> mice as Responder T cells and labeled with 5 μM CellTrace<sup>V</sup> Violet (CTV, ThermoFisher Scientific, MA, USA). CD3<sup>+</sup> MHC-II<sup>+</sup> splenocytes were sorted from C57Bl/6 CD45.2<sup>+</sup> mice and treated with 50 μg/mL of Mitomycin C (Calbiochem, MA, USA) to be used as antigen presenting cells (APC). Responder cells (5 × 10<sup>4</sup>) were polyclonally activated with 5 μg/mL anti-CD3 (clone 145-2C11, Biolegend) and Mitomycin-C (Mit-C) treated-APCs (1 × 10<sup>5</sup>), alone or co-cultured with different proportions of wt, het, or Nrp1KO Foxp3<sup>3GFP</sup>/Cre<sup>+</sup> Treg cells (1:1 to 1:4 Treg:ConcT cell) in round-bottom, 96-well plates with 200 μL RPMI, 10% FBS, 1% HEPES (Gibco, USA), 1% Penicillin/Streptomycin solution (ThermoFisher, USA) (complete RPMI or cRPMI) for 72 h. In some experiments, recombinant murine IL-10 (rmIL-10, Peprotech, USA) was added at final concentration...
of 10 ng/mL. Responder T cell proliferation was analyzed by CTW dilution by flow cytometry using FACSCanto II (BD Biosciences). Suppression was calculated as previously described (53). In brief, it uses the formula % Suppression = \( 1 - \frac{\text{DI}_{\text{Treg}}}{\text{DI}_{\text{Tresp}}} \times 100 \) % (where \( \text{DI}_{\text{Treg}} \) stands for the division index of responder cells with Treg cells, and \( \text{DI}_{\text{Tresp}} \) stands for the division index of responder cells activated without Tregs).

For transwell suppression experiments, CTV-stained Responder cells (\( 2.5 \times 10^6 \)) were stimulated with anti-CD3 (2 \( \mu \)g/mL) and Mitomycin-C (5 \( \mu \)g/mL) treated-APCs in 200 \( \mu \)L CRPMI in the bottom chamber of the multiwell plate, then 0.4 \( \mu \)m pore size transwell inserts (Corning, USA) were placed and \( w_t \), \( h_t \), or Nrp1KO Tregs (1.25 \( \times 10^5 \)) at different proportions (1:2 to 1:4 of Treg:convT cell ratio) activated with responder cells (1.25 \( \times 10^5 \)) and APCs (2.5 \( \times 10^5 \)) plus anti-CD3 in 100 \( \mu \)L on the top chamber for 72h. Proliferation of responder CD45.2+ convT cells was analyzed as previously mentioned.

**Ex vivo Suppression Assay**

CD45.1+ convT cells were sort-purified and i.v. injected into RAG-KO mice alone ["Group (i)"] or co-injected with CD45.2+Treg cells from \( w_t \) or Nrp1KO animals ["Group (ii)" and "(iii)", respectively], followed by allograft transplantation the next day as aforementioned. Fifteen-days post-surgery, dLN were harvested and pooled from mice of the same group. Cells were stained for viability ( ZombieDye NIR) and live CD45.1+CD4+ convT cells were FACS-sorted and co-cultured with freshly isolated CTW-stained CD45.2+CD4+ convT cells. T cells were activated with Mit-C-treated MHC-II+ APC plus soluble anti-CD3 (at 1:1 ratio of fresh CD45.2+convT:ex-vivo isolated CD45.1+convT cells). After 72h, the proliferation of CD45.2+CD4+ convTcell was analyzed by CTW dilution using flow cytometry as previously mentioned. Proliferation was calculated using the following formula: CD45.2+ convT proliferation = \( \frac{\text{DI}_{\text{fresh+ex-vivo}}}{\text{DI}_{\text{fresh}}} \times 100 \), where \( \text{DI}_{\text{fresh+ex-vivo}} \) stands for division index of fresh CD45.2+ convT cells activated with control \( w_t \) Tregs or ex vivo isolated CD45.1+ convT cells from RAG-KO animals, and \( \text{DI}_{\text{fresh}} \) stands for division index of fresh CD45.2+ convT cells activated alone.

**Western Blot**

Immunoblots were performed as previously described (54, 55). In brief, cell samples were lysed in RIPA buffer (10 mM Tris-HCl pH 6.8, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 0.1% SDS) on ice for 20 min. Lysates were vortexed and spun at 17,200 \( \times g \) for 20 min, and lystate supernatants were kept at \(-80^\circ C\) until further use. The protein concentration was determined using Pierce BCA Protein Assay Kit (Life Technologies, US). 5X reducing sample buffer (60 mM Tris-HCl pH 6.8, 25% glycerol, 2% SDS, 5% \( \beta \)-mercaptoethanol and 0.04% bromophenol blue) was added to lysates at proper proportion, samples were heated at 95°C for 10 min and cooled to room temperature before resolved by SDS-PAGE (SDS-polyacrylamide gel electrophoresis). Proteins were resolved under fully denaturing and reducing conditions, transferred to nitrocellulose membrane, blocked in 4% non-fat powdered milk in PBS-T (0.1% Tween-20) and probed with antibodies. The following antibodies were used for Immunoblot: anti-\( \alpha \)-Tubulin (clone B-7, Santa Cruz Biotechnologies, USA), and anti-Nrp1 (AF566, R&D Systems, USA), and peroxidase-coupled secondary antibodies anti-mouse IgG H+L (115-035-003, Jackson Immunoresearch, USA) and anti-goat IgG H+L (A27014, ThermoFisher, USA). Protein bands were detected using Pierce ECL Western Blotting Substrate (ThermoFisher, USA).

**Statistics**

Statistical analyses were performed using Student’s t-test, two-way ANOVA and log-Rank test using the software GraphPad Prism (CA, USA). Differences with \( p < 0.05 \) were considered significant. * \( p < 0.05 \); ** \( p < 0.01 \); *** \( p < 0.001 \); ns, not significant.

**ETHICS STATEMENT**

Mice were maintained under pathogen-free conditions at the animal facility located at Facultad de Medicina, Universidad de los Andes. All procedures were carried out according to the bioethics committee guidelines from Universidad de los Andes, and National Commission of Science and Technology (CONICYT).

**AUTHOR CONTRIBUTIONS**

MC-M performed experiments and analyze data. PC-K, FG-J, MR, CR, and AR participated in the execution of some experiments. OC discussed data and KP-L conceived and designed experiments, analyzed data, and wrote the manuscript.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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