Adoptive Transfer of Immunomodulatory M2 Macrophages Prevents Type 1 Diabetes in NOD Mice

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Macrophages are multifunctional immune cells that may either drive or modulate disease pathogenesis depending on their activation phenotype. Autoimmune type 1 diabetes (T1D) is a chronic proinflammatory condition characterized by unresolved destruction of pancreatic islets. Adoptive cell transfer of macrophages with immunosuppressive properties represents a novel immunotherapy for treatment of such chronic autoimmune diseases. We used a panel of cytokines and other stimuli to discern the most effective regimen for in vitro induction of immunosuppressive macrophages (M2r) and determined interleukin (IL)-4/IL-10/transforming growth factor-β (TGF-β) to be optimal. M2r cells expressed programmed cell death 1 ligand-2, fragment crystallizable region γ receptor IIb, IL-10, and TGF-β, had a potent deactivating effect on proinflammatory lipopolysaccharide/interferon-γ-stimulated macrophages, and significantly suppressed T-cell proliferation. Clinical therapeutic efficacy was assessed after adoptive transfer in NOD TID mice, and after a single transfer of M2r macrophages, >80% of treated NOD mice were protected against TID for at least 3 months, even when transfer was conducted just prior to clinical onset. Fluorescent imaging analyses revealed that adoptively transferred M2r macrophages specifically homed to the inflamed pancreas, promoting β-cell survival. We suggest that M2r macrophage therapy represents a novel intervention that stops ongoing autoimmune TID and may have relevance in a clinical setting. Diabetes 61:2881–2892, 2012

Macrophages have critical functions in both innate and adaptive immune responses. They are present in almost every tissue, recognize exogenous/endogenous danger signals through pattern-recognition receptors, produce various cytokines/chemokines that orchestrate immune responses at the site of inflammation, and function as professional antigen-presenting cells (APCs).

Two macrophage activation states have been defined in rodents and humans: “classically activated” (M1) cells have proinflammatory effector functions, and “alternatively activated” (M2) cells have anti-inflammatory properties. The existence of these different activation states implies prominent roles in different phases of an immunological response, i.e., inflammation versus resolution and tissue remodeling. M1 cells are identified by high expression of the enzyme inducible nitric oxide (NO) synthase, a potent respiratory burst, and secretion of proinflammatory cytokines such as tumor necrosis factor-α (TNF-α) and interleukin (IL)-12. The activation is induced by two signals, one toll-like receptor agonist, such as lipopolysaccharide (LPS), and one cytokine receptor-mediated signal, e.g., interferon-γ (IFN-γ) (1,2).

We now understand that there are subpopulations of M2 macrophages, different types of activation leading to different functional phenotypes. M2 cells are generally characterized by secretion of anti-inflammatory cytokines such as IL-10 and by low or no secretion of proinflammatory cytokines. IL-4 was initially identified as an inducer of M2 macrophages (3), and it was later discerned that IL-4, in combination with IL-13, enhanced induction of wound-healing macrophages (M2a) (4). Further investigations determined that stimulation with glucocorticoids (e.g., dexamethasone), IL-10, immune complexes in combination with toll-like receptor agonists, and transforming growth factor-β (TGF-β) could induce at least two further distinct M2 polarization states with immunoregulatory properties (M2b and M2c) (1,5). We have previously studied macrophage phenotypes from autoimmune-resistant and -susceptible rodent strains and have determined that there is a diversity of M1 activation phenotypes; autoimmune-susceptible strains have a common phenotype that contributes to prolongation of inflammation instead of its resolution (6). Similar genetically determined aberrant macrophage phenotypes have also been reported for NOD mice (7,8) and in human type 1 diabetes (T1D) (9,10). This indicates that there is a genetic predisposition for autoimmune susceptibility regarding macrophage phenotype and implies that individuals with autoimmune diseases may lack the ability to generate a cellular phenotype important in the resolution of inflammation.

Evidence of “immune regulatory” macrophage activity derives from cancer models in which tumor-associated macrophages have been reported to both suppress tumor immunity and promote tumor survival (8). Several in vitro (11,12) and in vivo disease studies (13–16) have investigated the regulatory role of macrophages in suppressing inflammation, the latter including models of multiple sclerosis, kidney disease, and spinal cord injury. Although these findings clearly indicate the important role of macrophages in the resolution of inflammation, there is no definitive consensus about their mechanism of action or optimal protocols for their induction.

TID is an autoimmune disease in which insulin-producing β-cells in the pancreas are attacked by leukocytes (macrophages and cluster of differentiation [CD]4+ and CD8+ T cells) (17), which leads to a subsequent loss of glucose control and acute complications. The most widely used

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animal model for investigation of T1D is the NOD mouse (18) in which diabetes spontaneously develops between 12 and 30 weeks of age after initial leukocyte infiltration into the pancreas (insulitis) between 3 and 5 weeks of age. Macrophages have been demonstrated to have a prominent role in T1D pathogenesis (19). Early M1 macrophage infiltration induces β-cell death and activates cytotoxic CD8⁺ T cells (20,21). Although many studies have provided convincing evidence of the destructive role of M1 macrophages in T1D, it was recently reported that transgenic NOD mice that did not spontaneously develop T1D expressed M2 macrophage–associated genes in the pancreas (22), and that the 10–20% of NOD mice that do not develop T1D possess protective macrophages with a phagocytic/immunosuppressive phenotype (23).

Previous studies have investigated the use of cell transfer as an immunomodulatory therapy in NOD mice, with most reports using highly suppressive regulatory T cells (Tregs) as an immunomodulatory therapy in NOD mice, with most reports using highly suppressive regulatory T cells (Tregs) (24,25). One drawback of this approach is that Treg antigen specificity is important for their suppressive ability (26), inferring that many Treg specificities might be required for the many pancreatic β-cell antigens implicated in T1D development. Adoptive cell transfer of tolerogenic dendritic cells has also been used in young NOD mice (5–8 weeks of age) during the early period of insulitis (27). For an optimal effect in a clinical setting, it would be advantageous to use cell therapy at the clinical debut of disease.

The role of M2 macrophages in T1D and adoptive transfer of regulatory M2 macrophages as a cell therapy in T1D have never been reported before. In this study, we elucidated the ability of IL-4/IL-10/TGF-β to induce an immunosuppressive M2 NOD macrophage phenotype (M2r). When M2r macrophages were adoptively transferred into late-stage prediabetic NOD mice, the onset of T1D was significantly reduced; advanced imaging analysis revealed protection of residual pancreatic islets concomitant with infiltration of the transferred M2r cells.

RESEARCH DESIGN AND METHODS

Animals. The female NOD/ShiLtJ (NOD) and NOD/ShiLtJ-TgFoxp3-EGFP/cre)1Jbs (NOD/Foxp3-GFP) mice (The Jackson Laboratory, Bar Harbor, ME) and NOD.Cg-Tg(TcrbBDC2.5, TcrbBDC2.5)1Dio/Doil (NOD-BDC2.5) mice used (from Dr. P. Hogquist, Karolinska Institutet, Sweden) were between 7 and 12 weeks of age. Diabetes development was assessed through blood glucose measurements (Freestyle mini glucose reader; Abbott, Abbott Park, IL), with blood glucose levels >14.1 mmol/L (250 mg/dl) for two consecutive days in fasted mice defining disease (Supplementary Fig. 1). All mice had normal fasting blood glucose levels <8 mmol/L, and experiments were approved by the local ethics committee.

Cell cultures. Macrophages (98% purity) were obtained as described previously (28) using macrophage colony-stimulating factor–conditioned and endotoxin-free Dulbecco’s modified Eagle’s medium supplemented with 20% FBS (complete medium). Splenocytes and lymphocytes from lymph nodes were cocultured in 96-well, flat-bottom plates or 0.4-µm transwell plates (Coming, Corning, NY) with 0.3 µg/ml anti-CD3 molecular complex (BD) stimulation of T lymphocytes. Fixation of APCs with 0.5% paraformaldehyde (PFA) has been described previously (29). Depletion of Tregs (99.5% effective) was conducted using magnetic microbeads (Miltenyi Biotec) and isolation of naive T cells using CD4+CD25- T-cell isolation kit II (Miltenyi Biotec). Proliferation of T cells was measured with [methyl-³H]thymidine (1 µCi/well; Amersham International, Amersham, U.K.). B2C2.5 T cells were stimulated with 500 µg/ml mitomoc 1040-31 peptide (Anaspec, Fremont, CA).

Reagents. Final concentrations were applied as follows: 50 ng/ml LPS, 200 nM dexamethasone (Sigma-Aldrich), 50 µg/ml vitamin D3 (Merck, Rahway, NJ), and 20 ng/ml recombinant mouse IL-4, IL-10, IL-13, and IFN-γ and recombinant human TGF-β1 (R&D Systems, Minneapolis, MN). Carboxyfluorescein succinimidyl ester (CSFE) (5–10 µM; Invitrogen) was used for measuring proliferation. Antibodies against CD86 (GLI), programmed cell death 1 ligand-1 (PD-L1) (MHI5), PD-L2 (TY25; all b Bioscience), CD3e (500A2; BD), CD3 (17A2; BioLegend, San Diego, CA), CD4 (GK1.5), CD8a (53-6.7), CD26L (ME-14; all Beckman Coulter), and CD44 (IM7; BD) and isotype control rat IgG2a (eBioscience) and IgG2b (eBioscience) were used for flow cytometric analyses. Dextran Alexa Fluor 467 (10,000 MW; Invitrogen) was used for endocytosis. Samples were run in a Gallios flow cytometer (Beckman Coulter, Brea, CA) and analyzed with Kaluza v1.1 (Beckman Coulter) and FlowJo v7.6.1 (Tree Star, Ashland, OR) software.

Cytokine analyses. ELISA kits were used for detection of secreted TNF, IL-6, TGF-β, and IL-10 in cell culture supernatants (Biologend and R&D Systems).

NO detection. NO activity was measured as nitrite in macrophage cell culture supernatants using the Griess reagent (Sigma-Aldrich) and measuring absorbance at 540 nm.

Tas pagan low-density arrays. RNA was extracted from stimulated macrophages (1 × 10⁶ cells/ml) using the RNeasy kit (Qiagen, Inc.), and RNA concentrations were measured by spectrophotometry (NanoDrop, Rockland, DE). cDNA was synthesized using SUPERSCRIPT II reverse transcriptase (Invitrogen). Data were analyzed using the comparative cycle threshold method and normalized using the geometric mean of two housekeeping genes (glyceraldehyde-3-phosphate dehydrogenase and hypoxanthine-guanine phosphoribosyltransferase). All experiments were related to untreated macrophage control.

Macrophage adoptive transfer. Macrophages stimulated with IL-4/IL-10/ TGF-β for 24 h or untreated macrophages (cultured in medium alone) were injected intraperitoneally with PBS as vehicle into groups of 8–12 mice at doses based on our previous studies (30).

Optical projection tomography. Dorsal pancreata from perfused and 4% PFA–fixed mice were dissected and frozen in PFA for 30–60 min at 4°C before stepwise transfer to 100% methanol and storage at −20°C. Staining for insulin and CD3 using whole-mount immunofluorescence and optical projection tomography scanning protocols were performed as previously described (31). The generation of tomographic reconstructions, images, and movies was performed as previously described (32) with the program NRecon v1.5.0 (SkyScan, Kontich, Belgium) visualization software module for Velocity v5.2.1 (Improvision; PerkinElmer, Waltham, MA).

Immunohistopathology. Islet cell–killed mice were perfused first with PBS and then fixed with 4% PFA, after which dorsal pancreata were isolated and prepared for cryosectioning. Transverse cryosections (14 µm) (Micron cryostat-microtome, Heidelberg, Germany) were mounted on glass slides and stored at −20°C. Staining comprised blocking of nonspecific binding with 5% goat serum/PBS/0.3% Triton X-100/0.01% sodium azide (Sigma-Aldrich) for 30 min, incubation with primary antibodies (guinea pig polyclonal anti-insulin; Abcam; rat anti-mouse CD3 molecule complex [1:50; BD]) overnight at 4°C and then for 1 h at room temperature with the respective goat anti–guinea pig Alexa Fluor 594 and goat anti-rat Alexa Fluor 488 (both 1:200; Molecular Probes) secondary antibodies. Sections were counterstained with DAPI and analyzed using a Leica DMIRE2 fluorescent microscope with DeltaPix software. Two to five slides per mouse were analyzed when insulin-positive islets were counted.

In vivo macrophage tracking. M2r macrophages and untreated macrophages were incubated for 5 min at 37°C with 3.5 µg/ml XenoLight D1(1,1′-dioctadecyl-3,3,3′,3′-tetramethylindocarbocyanine iodide Caliper Life Sciences, Hopkinton, MA), DiR (1,1′-dioctadecyl-3,3,3′,3′- tetramethylindocyanine perchlorate), or DiD (1,1′-dioctadecyl-3,3,3′,3′-tetramethylindocarbocyanine perchlorate); DiR is the dye for intraperitoneal injection. Mice were anesthetized and injected using IVIS spectrum (Caliper Life Sciences) with ex.710/em.760 filters for DiR. Images were analyzed with Living Image v3.2 (Caliper Life Sciences) software. Emission from control organs was subtracted from DiR cell–treated organs when quantification of macrophage migration was made. Retrieval of adoptively transferred macrophages from pancreata and pancreatic lymph nodes (PLNs) was conducted by enzymatic digestion with liberase, DNase (Roche), and hyaluronidase (Sigma-Aldrich). Debris and extracellular/endocrine cells were removed using M-Lympholyte (Cedarlane, Burlington, Ontario, Canada) phase separation.

Statistical analysis. Statistical significance was determined by Mann-Whitney U test (GraphPad software v5, San Diego, CA). A P value of <0.05 was considered significant, and error bars are presented in SEM.

RESULTS

Stimulation with IL-4, IL-10, and TGF-β induces a distinct M2r phenotype. We first screened a variety of stimulation protocols and characterized the induced M2 phenotypes. To assess the ability of different M2 stimuli to modulate the induction of proinflammatory M1 activation phenotypes, naive NOD macrophages were co-cultured with LPS/IFN-γ (M1) and each M2-inducing stimulus, alone or in different combinations. NO levels were significantly reduced by the synergistic effect of TGF-β and IL-10.
FIG. 1. IL-4, IL-10, and TGF-β induce a distinct anti-inflammatory M2 phenotype. Macrophages (1 × 10^5) were costimulated for 72 h with the respective stimulus and LPS/IFN-γ to determine levels of the proinflammatory mediators NO (A) and IL-6 (B). C: TNF measurement after activation with the respective stimulus for 24 h followed by secondary stimulation with LPS/IFN-γ for 24 h (48 h). D: Endocytosis assessed by stimulating 5 × 10^5 macrophages for 24 h with the respective stimulus followed by 4 h incubation with Alexa Fluor 647–coupled dextran. Bars represent mean fluorescence intensity (MFI). E: Levels of biologically active TGF-β secretion in 5 × 10^5 macrophages stimulated for 1 h, followed by 24 h incubation in fresh complete medium. F: IL-10 after costimulation of 5 × 10^5 macrophages for 1 h with the described stimulus and LPS, and further incubation in fresh complete medium for 24 h. G: Gene expression of costimulatory molecules identified using low-density arrays. Color patterns visualize fold gene expression relative to untreated controls (red indicates increase, green indicates decrease, and black designates no fold difference). H: PD-L1, PD-L2, and CD86 receptor expressions analyzed by flow cytometry (MFI) using receptor-specific antibody staining. Each color represents one macrophage stimulation (24 h) regimen. All results are representative of three independent experiments. Statistical comparisons were conducted against LPS/IFN-γ (A and B) or untreated (C–H) controls (black bars; n = 4). White bars represent negative controls. ND, not detectable. Error bars are presented in SEM. *P < 0.05.
IL-10 (Fig. 1A). Addition of IL-10 or TGF-β had significant downregulatory effects on IL-6 (Fig. 1B) and TNF (data not included) production compared with the LPS/IFN-γ control. The triple combination of IL-4/LI-10/TGF-β downregulated production of all three of these proinflammatory molecules.

To assess the fate of M2 phenotypes after secondary proinflammatory stimulation, macrophages were first M2 induced for 24 h, supernatants were washed away, and medium containing supplemented LPS/IFN-γ was added for another 24 h. Resultant proinflammatory TNF levels were significantly lower from the M2 cells initially activated by IL-4/LI-10/TGF-β (Fig. 1C).

Endocytosis is an important function during inflammation for clearance of both apoptotic cells and cellular debris from the local environment. The endocytic potential of induced M2 macrophages was assessed by uptake of fluorescently labeled dextran, and we determined that IL-4, either alone or in combination, induces enhanced endocytic activity in NOD macrophages (Fig. 1D).

As IL-10 and TGF-β both have deactivated effects on macrophages and are important during the resolution of inflammation (33–35), we next investigated if the M2 induction protocols led to their secretion by macrophages. Relatively high levels of biologically active TGF-β (Fig. 1E) were detected in macrophages that had been stimulated for only 1 h with TGF-β itself. The significant induction of IL-10 protein required TGF-β stimulation and concomitant activation with LPS (Fig. 1F), as previously reported (36). As LPS is proinflammatory, we also measured TNF in this setting and confirmed that the triple combination of IL-4/LI-10/TGF-β resulted in significantly lower levels of TNF (data not included).

Taken together, the findings of this screening of M2 induction protocols indicate that the triple combination of IL-4/LI-10/TGF-β induces a specific M2r macrophage phenotype that partly derives from synergistic effects of the three inducing cytokines. Each cytokine contributes with specific features such as endocytosis, phenotype stability, and secretion of TGF-β and IL-10.

**Differential expression of PD-L1/PD-L2 on M1/M2 macrophages.** A panel of B7 family (B7) costimulatory receptors, including CD80, CD86, B7-H3, B7-H4, V-set and Ig domain-containing 4 (Vsig4) (B7S1), PD-L1, and PD-L2, were screened for expression, as they have documented abilities to promote either activation or inhibition of T cells (37). Our results demonstrate that these receptors are differentially expressed at both transcription (Fig. 1G) and protein (Fig. 1H) levels, depending on the stimulation protocol. In particular, PD-L1 and PD-L2 were differentially expressed, mainly due to the actions of LPS/IFN-γ and IL-4, respectively. CD86 expression was strongly enhanced by LPS/IFN-γ, but flow cytometry data revealed that IL-4 also induces minor expression of CD86 and PD-L1 (Fig. 1H). There were no major expression differences evident for CD80 or Vsig4. Thus, the major finding was that elevated expression of PD-L1 in M1 macrophages and of PD-L2 in IL-4/LI-10/TGF-β–induced M2r macrophages distinguished these different activation phenotypes.

Additional gene transcription profiling revealed an M2 signature (high expression of Arg1, Ido1, Ido2, and Fcgri2b and low expression of Nos2, Th2, Ccl22, and Ccrl10) (Fig. 2). IL-10 did not induce any major gene expression by itself but had a synergistic effect on IL-4–stimulated macrophages. M2r macrophages retain an M2 signature after secondary proinflammatory stimulation both in vitro and in vivo. To further assess the plasticity of M2r macrophages, a set of experiments was performed (Fig. 3A).

Preactivated (24 h) M2r macrophages were washed and stimulated with LPS/IFN-γ for an additional 24 h before activation phenotyping (48 h). M2r macrophages significantly produced higher levels of IL-10 when stimulated with M1-inducing agents (Fig. 3B). PD-L2 expression was reduced after secondary stimulation but was still significantly higher relative to untreated (M0) and M1 control. CD86 was upregulated on M2r macrophages but still significantly lower than on M0 and M1 macrophages (Fig. 3C).

To further characterize the resultant phenotype of secondary–activated M2r macrophages, a set of gene targets was analyzed using PCR. M2–specific gene transcripts of Arg1, Tgbf1, Pdl2, C1163, Fcgri2b, Ido1, and Mrc1 were still present in secondary LPS/IFN-γ–stimulated M2r macrophages, but upregulation of M1–associated genes, such as Pdl1, Cds86, and Nos2, was also recorded (Fig. 3D). We additionally investigated the ex vivo phenotype of DiD–stained M2r macrophages in PLNs and pancreata from NOD-BDC2.5 mice 1 week after transfer (Fig. 3E). Around 78% of the DiD+ M2r macrophages expressed PD-L2 before transfer, and ex vivo analysis of retrieved DiD+CD11b+ M2r macrophages revealed that >50% still expressed PD-L2, although the majority of the transferred cells also coexpressed CD86 in the PLN. A similar scenario was evident in the pancreas but fewer M2r macrophages expressed PD-L2 (~42% in total). In summary, these data indicate that transferred M2r macrophages retain an M2 signature and do not adopt a dominant M1 phenotype on access of the inflamed pancreas.

**Adoptive cell transfer of M2r protects mice from T1D.** The cumulative in vitro data indicated that IL-4/LI-10/TGF-β induced an optimal suppressive M2r macrophage phenotype. The in vivo immunoregulatory activity of these cells in a setting of autoimmune disease was next investigated through their adoptive intraperitoneal transfer into 16-week-old, prediabetic, NOD mice, a time point just prior to clinical onset. Strikingly, even at this late time of intervention, 83% of the M2r–treated mice (single treatment) were protected from T1D, whereas only 25% of mice injected with M0 macrophages or vehicle (PBS) were diabetes free after 3 months of follow-up (Fig. 4A). This result indicates the potent ability of M2r macrophages to halt an ongoing autoimmune disease process.

**M2r–treated mice exhibit preserved β-cell survival.** Given the impressive clinical effect of M2r therapy, the pancreata from M2r and M0 macrophage–treated mice were examined for endocrine and immune activities. After dissection at different time points, three-dimensional reconstructions of pancreata were performed using optical projection tomography (Fig. 4B and Supplementary Videos 1–5). Control mice at 16 weeks of age clearly had very active insulitis and low numbers of insulin–positive islets (compared with 5-week-old NOD mice in which insulin–positive islets were abundant) (38). Conversely, a higher number of insulin–producing β-cells were detected in mice treated with M2r macrophages, underlining that transfer of M2r macrophages led to diminished β-cell destruction (Fig. 4B). Immunohistochemical analyses of pancreata revealed similar findings (Fig. 4C), with a modest, yet significant, quantifiable difference in pancreatic islet numbers in M2r–treated animals (Fig. 4D).

Macrophages predominantly migrate to the pancreas/PLN. Live cell tracking using the stable cell tracer DiR was used to study the fate of adoptively transferred macrophages.
Macrophage migration was already evident 2 h postinjection; the cells were clustering near the pancreas and spleen (Fig. 5A), with no major differences between analyses on days 3, 6, and 8 postinjection, respectively. To confirm specificity, organs were dissected out from mice on day 3 and analyzed individually (Fig. 5B). A significant majority of the fluorescent emissions from both M0 and M2r macrophages were localized to the pancreas and the PLN (Fig. 5C). Additional immunohistochemical analyses confirmed the presence of fluorescent M2r macrophages in close proximity to the remaining islets, with colocalization of CD11b immunostaining (day 3 depicted in Fig. 5D).

Taking these in vivo data together, we conclude that adoptively transferred M2r macrophages migrate to the inflamed pancreas in NOD mice, resulting in protection from overt clinical T1D onset.

**M2r suppress T-cell proliferation in vivo.** To address the local in vivo mechanism of action of transferred M2r macrophages, NOD-BDC2.5 mice were used because the majority of their T cells recognize β-cell antigen. NOD-BDC2.5 mice were treated with M2r or M0 macrophages, PLNs were dissected out 1 week posttreatment, and antigen-specific proliferation was assessed in vitro. Lymphocytes from M2r-treated BDC2.5 mice proliferated less (16518 ± 3,333 counts per min [CPM]) than lymphocytes from M0-treated mice (32807 ± 7,911 CPM) upon restimulation with β-cell antigen peptide, clearly indicating immunomodulation of T cells by M2r macrophages in vivo (Fig. 6A). The relative numbers (percentage of cells and absolute numbers) of PLN CD4+ and CD8+ T-cell subsets were not affected by

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**FIG. 2.** IL-4 dominantly regulates gene expression in 4 h–stimulated macrophages. Color patterns visualize fold gene expression relative to untreated control. Red color indicates an increased fold expression and green color indicates a decreased fold expression. Black color designates no fold difference. Representative data from three individual animals.
M2r treatment (Fig. 6B). There was a trend ($P = 0.06$) of altered activation state of these CD4+ T cells in M2r-treated mice, as determined by CD62L and CD44 expression (Fig. 6C). Similar analyses of cells recovered from the pancreata of M2r-treated mice also revealed no differences in T-cell subset compositions, although a trend ($P = 0.06$) of lower CD8+ T-cell activation was observed (Fig. 6D). These experiments were repeated in NOD-FoxP3-GFP mice. Again, there were no differences in the number of T-cell subsets or Tregs in PLNs from M2r-treated mice (Fig. 6E).
FIG. 4. M2r macrophages protect NOD mice from T1D by protecting pancreatic β-cells. A: IL-4/IL-10/TGF-β–stimulated macrophages (2.5 × 10^6) (M2r, blue, n = 12), untreated macrophages (M0, red, n = 8), or vehicle (PBS, black, n = 8) were intraperitoneally injected into 16-week-old prediabetic NOD mice (arrow). The M2r-treated group was significantly protected compared with M0 and PBS groups, as independently statistically analyzed using the Mantel-Cox test in a Kaplan-Meier survival graph. These results are representative data from two independent experiments with a similar outcome. B: Organs were stained with anti-insulin (red) and anti-CD3 (green)–specific antibodies prior to three-dimensional reconstruction using optical projection tomography. Yellow represents colocalization of CD3 and insulin staining, indicating insulitis. Organs from 16-week-old mice represent three individual animals; 21- and 26-week-old organs represent two individual animals. C: After M2r or M0 macrophage transfer, pancreata were dissected 8 weeks later (24 weeks of age), and cryosections were stained with anti-insulin (red) and anti-CD3 (green) (n = 5). D: Insulin+ islets were counted manually from the sections in two to five transverse sections per animal to obtain the average number of insulin+ islets per section (n = 4). *P < 0.05. (A high-quality digital representation of this figure is available in the online issue.)
FIG. 5. Macrophages predominantly migrate to the pancreas and PLN. A: NOD mice (10–12 weeks of age) received 3 × 10^6 DiR-labeled M2r or M0 macrophages intraperitoneally. Mice were anesthetized prior to anterior imaging after 2 h and 1, 3, 6, and 8 days post–macrophage injection. Bright yellow or dark red represents high or low photon counts, respectively, and white arrows indicate injection sites. Two individual mice were analyzed, and the data represent two individual experiments. B: After transfer, the liver, kidney, spleen, pancreas, and PLNs were dissected after 3 days post–macrophage injection. The image is representative of eight individual mice (four treated with M0 and four with M2r). C: Photon count from each organ in B was quantified, and control organ photon count was subtracted to obtain the real macrophage emission. Error bars are presented in SEM. D: NOD mice received 2 × 10^7 Dil-labeled M2r macrophages (red) intraperitoneally, and pancreata were dissected at day 3 and stained for CD11b (green) and DAPI (blue). The data are representative of four different mice. (A high-quality digital representation of this figure is available in the online issue.)
FIG. 6. M2r macrophages suppress ex vivo T-cell activity. M2r or M0 macrophages (2.5 × 10^6) were intraperitoneally injected into 12–16-week-old NOD-BDC2.5 mice. A: PLNs were dissected after 1 week, and lymphocytes were restimulated with BDC2.5 mimotope for 72 h to induce proliferation. Readout was CPM. B: PLNs were dissected after 1 week, and lymphocyte subsets were analyzed by flow cytometry for cell numbers (percent and absolute values). C: Activation status of CD4^+ subset assessed by CD44 and CD62L expression. D: Pancreata were dissected after 1 week, and lymphocyte subsets were analyzed by flow cytometry for cell numbers (percent and absolute values). E: M2r macrophages (2–3 × 10^6) or control (PBS) were intraperitoneally injected into 12–13-week-old NOD-FoxP3-GFP mice. PLNs were dissected after 1 week, and T-cell subset numbers were analyzed. The data from A–D represent two independent experiments (n = 4). The data in E represent pooled data from four independent experiments. Error bars are presented in SEM. *P < 0.05.
FIG. 7. TGF-β is important for M2 macrophages to suppress T-cell proliferation and the induction of Tregs in vitro. A: Suppression of T-cell proliferation (CPM) by stimulating 25 × 10^3 macrophages for 24 h and further coculture with splenocytes and αCD3 for 72 h. B: Macrophages (25 × 10^3) were cultured and stimulated for 24 h in a transwell upper chamber. Splenocytes and αCD3 were added to the lower chamber, and cells were coincubated for 72 h and assessed for proliferation (CPM). C: Cell contact-dependent suppression of proliferation (CPM) evaluated by stimulating 25 × 10^3 macrophages for 24 h followed by fixation and coculture with T cells, splenocytes, and αCD3 for another 72 h. D: CFSE-stained splenocytes cocultured with 25 × 10^3 pre-stimulated macrophages, as described above, for 96 h. Cells were either gated for CD4+ or CD8- (left), and CFSE dilution was analyzed (right). E: Macrophages (25 × 10^3) were cultured and stimulated for 24 h before coculture with splenocytes from NOD-FoxP3-GFP mice for 96 h. Cells were gated for CD4+ cells, and the percentage of GFP (FoxP3)-expressing cells was analyzed. F: Macrophages (25 × 10^3) were cultured with CD25-depleted splenocytes from NOD-FoxP3-GFP mice for 96 h. Cells were gated for CD4+ cells, and the percentage of CD25+GFP+ expressing cells was analyzed. G: Macrophages (25 × 10^3) were cultured with CD4+CD62L+CD25- T cells from NOD mice for 96 h. Cells were gated for CD4+ cells, and the percentage of
nor were there major changes in T- or B-cell numbers detected in the spleen or other lymph nodes of treated NOD-FoxP3-GFP mice (data not included).

**TGF-β has a major role in the suppressive and Treg induction activities of M2r macrophages.** Macrophage regulation of T-cell activities has been reported during the recent decade in both tumor and inflammation biology settings, but the nature of the cytokines that induce these suppressive abilities in macrophages is poorly understood. To confirm the in vivo findings of the suppressive action of M2r macrophages, an in vitro suppression assay was used, using anti-CD3 splenocyte activation. Optimization of cell densities led to the 1:16 (macrophage to T cell) ratio being selected for use in all further suppression assays (data not included). M2 induction by TGF-β, either alone or in combination with IL-4/IL-10, was crucial for macrophage suppression of splenocyte proliferation (Fig. 7A). Both a transwell system and macrophage fixation were used in suppression assays to assess the relative contributions of cell contact–independent and –dependent suppressive mechanisms, respectively. TGF-β-stimulated macrophages suppressed T-cell proliferation both in the transwell (Fig. 7B) and after fixation (Fig. 7C). Taken together, these results indicate that TGF-β has an important role in the regulatory phenotype of macrophages to suppress T-cell activation mediated both by cell contact and via secreted factors. NOD splenocytes were also stained with carboxyfluorescein diacetate succinimidyl ester (CFSE) and cocultured with induced M2r macrophages, demonstrating that proliferation of both CD4+ and CD8+ T-cell subsets is suppressed (Fig. 7D).

When combined with IL-4/IL-10, TGF-β–treated macrophages suppressed T-cell proliferation, secreted TGF-β, and expressed PD-L2. As these represent molecules associated with Treg induction, the ability of M2 cells to induce Tregs was assessed in vitro. TGF-β pretreatment of macrophages led to an increase in the percentage of Tregs induced from naive T cells in vitro (Fig. 7E). Interestingly, only TGF-β/IL-10–stimulated macrophages had the ability to significantly induce Tregs from CD25−depleted splenocytes (Fig. 7F). We additionally used naive CD4+ T cells (CD62L+CD25−) to confirm that M2-stimulated macrophages induced Tregs (Fig. 7G).

**DISCUSSION**

Macrophages are important in the pathogenesis of many autoimmune diseases, being the first cells to infiltrate the pancreata of NOD mice (39), major effector cells in experimental multiple sclerosis (40), and major TNF producers in rheumatoid arthritis (41). In contrast to the tissue-destructive functions of M1 macrophages in autoimmune diseases, the immunosuppressive and tolerogenic ability of M2 macrophages and dendritic cells has stimulated an emerging interest for these myeloid cells and their immunomodulatory roles in regulation of inflammation. The major finding of our current study is that a single adoptive transfer of in vitro induced immunosuppressive M2r macrophages can prevent imminent development of clinical T1D in NOD mice, illustrating the potential of this myeloid cell therapy to counteract an aggressive pathological process.

In screening to define an optimal suppressive macrophage activation phenotype in vitro, we applied an extensive panel of activation protocols encompassing reported M2r-inducing protocols. Taken together, these in vitro phenotypic analyses revealed that a spectrum of M2 phenotypes was induced via different induction protocols and that different readouts were required to distinguish these individual phenotypes. We finally selected the novel combination of IL-4/IL-10/TGF-β for further study, a cytokine cocktail encompassing both M2a and M2c phenotypes (5) and indicating both potential regulatory and wound-healing functions of the induced M2r phenotype. Although PD-L1 and PD-L2 receptors have been discordantly reported as being either inhibitory of (42–44) or activating (45) coreceptors depending on the inflammatory scenario, we determined the expression signature of PD-L1low/PD-L2high/CD86low in M2r cells, which contrasted with PD-L1high/PD-L2high/CD86high expression in M1 cells.

The individual effects of IL-4, IL-10, and TGF-β in the M2r induction protocol either synergized or antagonized expression of specific molecules, but this activation phenotype was functionally suppressive via both secreted and surface-expressed molecules. The production of both TGF-β and IL-10 and low levels of TNF after co- or secondary stimulation with LPS clearly indicates their potential regulatory role. Functional analyses revealed that this novel anti-inflammatory phenotype could not only suppress T-cell proliferation in vitro but also modulate the ex vivo activation of PLN T cells. Considering that transferred M2r macrophages migrated to the inflamed pancreas, the deactivation of infiltrated proinflammatory macrophages, dendritic cells, or autoreactive T cells within the target organ could be one of the mechanisms explaining the clinical efficacy in preventing T1D development posttransfer into NOD mice, while they might also induce wound-healing processes in the target organ. Once in the proinflammatory environment of the pancreas, the transferred cells retained an M2 signature. The anti-inflammatory action of the M2r macrophages thus appears to be central to the clinical therapeutic effect, although the potential contribution of local in vivo Treg modulation cannot be ruled out.

We previously demonstrated an APC transfer therapy in a setting of experimental neuroinflammation (30) by adoptively transferring splenic major histocompatibility class II (MHCII+) cells into mice with previously induced encephalomyelitis and demonstrating a significant therapeutic effect of this cell therapy. During recent years, this has become an intense area of research, and other research groups have confirmed the therapeutic concept in both autoimmune neurologic and renal disease settings, indicating that the approach may be used in a variety of inflammatory settings. That T1D could be prevented by so late an intervention in NOD mice is striking and indicates both the potent immunosuppressive activity of the M2r phenotype and their novel role in T1D. Based on one injection, and in a nonantigen-dependent fashion, this cell therapy is clinically an attractive option for combining immunosuppressive and wound-healing activities.

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CD25+FoxP3+ expressing cells was analyzed. The results are representative of three independent (A–F) experiments and one experiment (G). Statistical comparison was conducted against untreated macrophage control (black bars; n = 4). Control represents splenocytes without macrophages (white bars). Error bars are presented in SEM. *P < 0.05.
No potential conflicts of interest relevant to this article were reported.

R.P. designed the study, wrote the manuscript, and performed research. P.A., A.G., S.Mi., X.-M.Z., and S.Ma. performed research. D.H. designed the study and wrote the manuscript. R.A.H. designed and supervised the study and wrote the manuscript. All authors analyzed data and revised the manuscript. R.A.H. is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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2892 DIABETES, VOL. 61, NOVEMBER 2012 diabetes.diabetesjournals.org