The Hepatocyte growth factor (HGF) and its receptor (MET) promote several physiological activities such as tissue regeneration and protection from cell injury of epithelial, endothelial, neuronal and muscle cells. The therapeutic potential of MET activation has been scrutinized in the treatment of acute tissue injury, chronic inflammation, such as renal fibrosis and multiple sclerosis (MS), cardiovascular and neurodegenerative diseases. On the other hand, the HGF-MET signaling pathway may be caught by cancer cells and turned to work for invasion, metastasis, and drug resistance in the tumor microenvironment. Here, we engineered a recombinant antibody (RDO24) and two derived fragments, binding the extracellular domain (ECD) of the MET protein. The antibody binds with high affinity (8 nM) to MET ECD and does not cross-react with the closely related receptors RON nor with Semaphorin 4D. Deletion mapping studies and computational modeling show that RDO24 binds to the structure bent on the Plexin-Semaphorin-Integrin (PSI) domain, implicating the PSI domain in its binding to MET. The intact RDO24 antibody and the bivalent Fab2, but not the monovalent Fab induce MET auto-phosphorylation, mimicking the mechanism of action of HGF that activates the receptor by dimerization. Accordingly, the bivalent recombinant molecules induce HGF biological responses, such as cell migration and wound healing, behaving as MET agonists of therapeutic interest in regenerative medicine.

In vivo administration of RDO24 in the murine model of MS, represented by experimental autoimmune encephalomyelitis (EAE), delays the EAE onset, mitigates the early clinical symptoms, and reduces inflammatory infiltrates. Altogether, these results suggest that engineered RDO24 antibody may be beneficial in multiple sclerosis and possibly other types of inflammatory disorders.

Keywords: HGF, MET, PSI domains, antibody engineering, regenerative medicine, anti-cancer therapy, multiple sclerosis, experimental autoimmune encephalomyelitis
INTRODUCTION

Hepatocyte growth factor (HGF), also known as scatter factor, is a multi-functional cytokine that was originally cloned as a potent mitogen for hepatocytes in primary cultures (1) and was later identified as a cell motility factor for epithelial, muscle and nerve cells (2). HGF is produced and secreted as an inactive pro-HGF form (92 kDa) by stroma cells such as macrophages and fibroblasts. Secreted pro-HGF is cleaved between Arg494 and Val495 to obtain active HGF, which is constituted by two chains, covalently linked by a disulfide bridge: a heavy chain of 62 kDa and a light-chain of 32-36 kDa. The activation of HGF is mediated by various proteases such as urokinase, plasminogen activator, and HGF activator (3, 4). The latter is the most efficient and is itself activated upon thrombin cleavage of its precursor (5). HGF is the ligand of the MET receptor tyrosine kinase (6, 7). The biologically active αβ heterodimeric HGF contains a high affinity MET-binding site in the α chain (8) and a low-affinity MET-binding site in the β chain (9). Only the active form of HGF is able to elicit MET-mediated biological activities.

MET receptor is synthesized as a single-chain precursor that is cleaved into two disulfide bond-linked α and β subunits of 50 and 140 kDa, respectively, by the furin protease in the Golgi apparatus (10). The αβ heterodimer consists of an N terminal extracellular domain (MET ECD), a trans-membrane domain, a juxta membrane region, an intracellular tyrosine kinase domain and a C-terminal tail. The MET ECD is composed of a Semaphorin (SEMA) domain, a Plexin-Semaphorin-Integrin domain (PSI), and four Immunoglobulin-like regions found in Plexins and Transcription factor (IPT1-4). The SEMA domain is necessary and sufficient for HGF binding (11) and is required for receptor dimerization and activation (12). The IPT3,4 regions have also been proposed as a further MET binding site for the HGF α-chain (13). HGF binding upon MET activates the receptor leading to several biological responses including cell scattering, motility, survival and differentiation. MET is primarily expressed in epithelial and endothelial cells (14), but also in some myoblasts and neuronal precursors thus contributing to the development of muscular and nervous structures (15–18). The normal expression and function of MET and HGF are fundamental during embryogenesis, promoting growth and development of hepatocytes, placental trophoblasts and myoblasts (16, 18, 19). Furthermore, MET activation is involved in organ growth and after injury regeneration, angiogenesis, wound healing, scattering, and proliferation (20–22).

Thus, MET activation is an interesting target in regenerative medicine and HGF role has been investigated in a panel of injury/disease models (23). The use of MET-deleted mice demonstrated the involvement of HGF and MET in regeneration, protection, and homeostasis of tissues. Many studies demonstrated that HGF is a powerful neurotrophic factor in the nervous system with beneficial and protective effects in various animal disease models. Thus, a possible therapeutic application of HGF is suggested for the treatment of neurological, neurodegenerative, and psychiatric disorders (see Desole et al., 2021 for a recent review) (24). HGF and MET play also important cardioprotective roles in the injured heart, by promoting pro-survival effects in cardiomyocytes, such as protection from apoptosis, autophagy, and genotoxicity (25, 26). Moreover, HGF drives migration and proliferation of cardiac stem cells (27, 28). Furthermore, studies on MET-deleted mice revealed a fundamental inhibitory role of HGF in the progression of chronic inflammation and fibrosis. Chronic tissue injury and inflammation have been associated with the onset of fibrosis. HGF treatment resulted in reduced fibrosis and improved tissue functions in a panel of disease models, such as liver cirrhosis, chronic kidney disease, and chronic fibrosis, due to its anti-apoptotic and anti-inflammatory activity (29–37). In the experimental autoimmune encephalomyelitis (EAE) model of multiple sclerosis (MS), mesenchymal stem cells transplantation ameliorated EAE clinical symptoms and this benefit is caused in large part by the production of HGF (38).

On the other hand, HGF-MET signaling plays a role in cancer cells, conferring malignant features such as invasion, metastasis, and drug resistance in the tumor microenvironment (22). The oncogenic gain of function of MET is obtained through different mechanisms. Genetic alterations of MET gene, such as gene amplification, point mutations or transcriptional activation, result in constitutive activation of MET kinase, which turns into a powerful oncogenic receptor insensitive to ligand regulation (39). Targeting of constitutively active MET by means of inhibitors is useful to blunt tumor growth in experimental models and in patients (40, 41). Moreover, stressful signals that promote transcriptional activation of MET in cancer cells may also induce HGF upregulation in tumor stromal cells (42), feeding a positive stimulatory circuit of HGF-MET signaling that allows cancer cells to convert the anti-apoptotic and pro-migratory activities typically used in tissue regeneration and repair into pro-invasive and pro-metastatic behaviors (43). Thus, several MET-targeting agents, including HGF and MET antibodies, as well as small molecule kinase inhibitors, are currently envisaged as anti-cancer therapeutics (44).

In a previous work, we generated a mouse monoclonal antibody (mAb), known in the art as DO24 (45), endowed with MET agonist activity (46). We also demonstrated that this agonist mAb protects cardiomyocytes from hypoxic and chemotherapy injury, thus displaying potential therapeutic
properties (25, 26). Here, we describe the generation and characterization of a synthetic antibody and antibody fragments derived from DO24 mAb. We recovered and cloned the VH (heavy chain variable domain) and VL (light chain variable domain) cDNA sequences from DO24 mouse hybridoma into appropriate vectors, introduced the vectors into host mammalian cells, and achieved expression of adequate amounts of functional antibody. The recombinant antibody (RDO24_mIgG2a) and its fragments (RDO24_mFab2 and mFab) were thus generated in vitro and assessed for MET agonism. Only the bivalent molecules demonstrated agonistic activity. Moreover, binding of recombinant antibody required the PSI and not the SEMA domain of MET, suggesting different molecular mechanisms for receptor activation. The stable full-size RDO24 was tested in the EAE model and showed a delay in the disease onset and severity. These engineered anti-MET molecules are promising candidates for therapeutic purposes and will be useful tools for further exploring MET receptor activation.

MATERIALS AND METHODS

Design, Synthesis, and Purification of the RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab

Molecules have been designed by inserting the synthetic codon optimized VH and VL cDNA sequences from DO24 mouse mAb in pcDNA3.1 vector which carry the murine constant IgG2a (immunoglobulin 2a) heavy chain (CH1, CH2, CH3) and the murine constant k light chain (CL), respectively. DNA plasmids were produced as endotoxin-free preparations and sequence-verified to confirm identity prior to mammalian ExpiCHO-S transient transfection (FlowEighteen38, Porto, PT). RDO24_mIgG2a proteins were purified using a HiTrapMabSelect Sure Protein A 5 mL column (GE Healthcare, Buckinghamshire, UK) on an ÄKTA Pure 25L FPLC system. RDO24_mFab2 and RDO24_mFab proteins were purified using a HisTrap HP 5 mL column (GE Healthcare). Analysis of the purified molecules was performed by SDS-PAGE under reducing and non-reducing conditions, followed by GelCode Blue Stain reagent (Pierce, Waltham, MA). The production of DO24 mAb, the antibody from which RDO24 molecules were derived, was performed as previously described (46). Briefly, DO24 was produced from hybridomas obtained with the fusion between the immune spleen cells from Balb/c mice (Charles River Laboratories, Wilmington, MA, USA) immunized with GTL-16 cells, where the MET gene is amplified and overexpressed, and the P3.X63.Ag8.653 myeloma cells.

Cell Culture and Materials

H9c2, A549, HEK293T, GTL16, TOV-112D, MDCK and HPAF-II cell lines were purchased from the American Type Culture Collection (ATCC, Manassas, VA, USA). Rhabdomyosarcoma cells were kindly provided by Riccardo Taulli (University of Turin). HUVECs were obtained from 15 samples following parental consent, and grown in gelatin-coated plates in their own complete medium (M199 medium supplemented with 10% FCS, 0.02% extract of bovine brain, and 0.015% porcine heparin). A549 and GTL16 cells were cultured in RPMI medium; H9c2 and rhabdomyosarcoma cells were grown in DMEM medium, HEK293T were cultured in Iscove’s medium, and MDCK and HPAF-II in EMEM medium. TOV-112D were cultured using a 1:1 mixture of MCDB 105 medium and medium 199 supplemented with 15% fetal bovine serum (FBS). RPMI, Iscove, DMEM, and EMEM media were supplemented with 10% FBS, 1% penicillin, 1% streptomycin and 1% L-Glutamine. Cells were incubated under 5% CO2 at 37°C, were passed regularly and sub-cultured to ~80/90% of confluence. Unless specified, all materials were from SigmaAldrich (St. Louis, Missouri, USA). Anti-pMET (Y1234/1235; 3077), -MET (D1C2; 8198), -ERK (T202/Y204; 4376), -pAKT (S473; 9271), -pCREB(S133; 9198), and -CREB (4820) antibodies were purchased from Cell Signaling Technology (Danvers, MA, USA), and anti-ERK (C14; SC-154) and -AKT (C-20; SC-1618) from Santa Cruz (Santa Cruz, CA, USA). HGF (Recombinant Human Hepatocyte Growth Factor NS0-expressed) was purchased from R&D systems (Minneapolis, Minnesota, USA).

Surface Plasmon Resonance

The kinetic constants of RDO24, Fab2, and Fab with recombinant human MET ECD-Fc (R&D) were measured using a Biacore T100 instrument (GE Healthcare) and the CM5 chip, following standard procedures. Human MET ECD-Fc (pH 4) was immobilized onto the surface of a single channel of the CM5 sensor chip by amine coupling. In a single-cycle kinetics experiment, RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab solutions were separately injected in four flushes at increasing concentrations (from 41 nM to 330 nM for RDO24_mIgG2a or from 62 nM to 500 nM for RDO24_mFab and RDO24_mFab2) over the human MET ECD-Fc functionalized sensor surface, using HBS-EP+ (Cytiva) as running buffer, with a contact time of 120 seconds. The reference flow cell was used as a control surface for refractive index change and nonspecific binding. A long dissociation phase (600 seconds) and a single regeneration step followed the last sample injection (NaOH 50 mM, 30 seconds of contact time), without regeneration between each sample injections.

Flow Cytometry

Human GTL16, A549, HEK293T, HUVEC, rat H9c2, and mouse rhabdomyosarcoma cells were resuspended in PBS 1% FBS and then stained for 15 min at room temperature, in the dark, with RDO24_mIgG2a labeled by PE-Cy7 fluorochrome, using Lightning-Link (Innova Biosciences, Cambridge, UK). To exclude died cells the stained cells were resuspended in Phosphate Buffered Saline (PBS) Dapi 0.2X solution. The negative control was the unstained cells. Samples were analyzed on a CyAn™ ADP LX nine-color analyzer (Beckman Coulter, Brea, CA, USA).
**Lentiviral Vectors and Engineering of MET Wild-Type and MET Deletion Mutants**

MET transmembrane receptors described in this work have been generated by standard PCR and genetic engineering techniques. All of the proteins conserve the signal peptide of their parental polypeptide at the N-terminus. The receptors are identical to MET wild-type (WT: total amino-acid length 25-1390), except for the deletion of SEMA domain (MET ΔSEMA: Δ25-516) or SEMA and PSI domains (MET ΔSEMA-PSI: Δ25-562) or IPT domains (MET ΔIPT: Δ563-932) or PSI and IPT domains (MET ΔPSI-IPT: Δ517-932). The cDNAs encoding all of the engineered proteins were subcloned into the lentiviral transfer vector pRRL2. Vector stocks were produced as previously described (47).

**Immunoprecipitation Assay**

TOV-112D cells without or with ectopic expression of MET WT or MET deletion mutants were lysed with cold RIPA buffer in the presence of 1 mM Na3VO4 and a cocktail of protease inhibitors (all from Sigma-Aldrich). Total protein lysates were incubated at 4°C overnight on rotor with RDO24 mAb, then Sepharose- protein A (GE Healthcare) was added and the samples were incubated for additional 2 h at 4°C. As control, an equal amount of total proteins was incubated with Sepharose protein A in the absence of antibodies. After five washes with cold RIPA buffer, immunoprecipitated proteins were eluted with boiling Laemly buffer and analyzed by western blotting.

**Enzyme-Linked Immunosorbent Assays**

To analyze the specificity and selectivity of RDO24_mIgG2a, 96well EIA/RIA plates (Costar, #3590, Corning, NY, USA) were coated at 4°C overnight with 100ng/well recombinant Fc-fused extracellular domain-fragment (ECD-Fc) of human MET (Recombinant Human HGFR/c-MET Fc Chimera His-tag Protein, R&D) or RON ECD (Recombinant Human MSP R/RON, R&D) or SEMA4D ECD-Fc (Recombinant Human Semaphorin4D Fc Chimera Protein, R&D). Following three washes with PBS with 0.05% Tween 20, they were blocked with PBS-0.5% BSA (bovine serum albumin) for 1 h at 37°C. Wells were then washed thrice with PBS-0.05% Tween and then RDO24_mIgG2a or anti-RON (RON monoclonal antibody, MA5-31073, Invitrogen) or antiSEMA4D (SEMA4D monoclonal antibody, eBio133-IC6, Invitrogen) were incubated at increasing concentration (0-100 nM) at 4°C overnight. After three washes with PBS with 0.05% Tween 20, wells were incubated with 1:5000 mouse horseradish peroxidase-conjugated secondary antibody (Jackson Laboratory, Bar Harbor, ME, USA) for 1 h at room temperature, followed by three more washes with PBS with 0.05% Tween 20. Tetramethylbenzidine (TMB) substrate solution (100 µl/well) was then added and quenched after 3-4 minutes with 25 µl of 2 M H2SO4. Colorimetric assay was quantified by the multi-label plate reader VICTOR-X4 (Perkin Elmer Instruments INC.). Binding data were analyzed and fitted using Prism software (Graph Pad Software, San Diego, CA, USA).

**Scatter and XCelligence Assays**

For end-point analysis, MDCK cells (12000 cells/well) were seeded in 96-well plates in complete culture medium. After 24 h, cells were incubated in the presence of RDO24_mIgG2a, mFab2 and mFab (50 nM), or HGF (0.5 nM) for 20 h.
Cells were fixed with 11% glutaraldehyde and stained with 0.1% Crystal Violet (SigmaAldrich). For real-time cell motility assay, HPAF-II cells (10000 cells/well) were seeded in E-plates (Roche Diagnostics, Mannheim, Germany) in complete culture medium and treated as above. Electrical impedance was monitored continuously for 24 h using an X-Celligence RTCA device (Roche Diagnostic). The electronic readout of cell-sensor impedance is displayed in real-time as cell index, a value directly influenced by cell shape and spreading. The induction of cell flattening and cell dissociation results in an increase of the cell index.

Wound Healing Assay

H9c2 cardiomyoblasts (150000/well) were plated in 24-well plates and maintained in DMEM 10% FBS until confluence, and then were incubated in DMEM 0.5% FBS for 18 h. A scratch wound was made by scratching with a 10-µL pipette tip across the center of the well. Then cells were washed with PBS, left untreated or incubated for 24 h with HGF, whole RDO24 mAb, and fragments. Images of wound at the start moment and after the treatment were taken with DMRI Leica inverted microscope. Migration was quantified by evaluating the area of wound at time zero (A0) and at time after the treatment (Ay = 24 h). Normalization and quantification on the basis of three independent experiments were obtained by the formula (A0–Ay)/A0.

Animals

All experimental procedures were carried out at Neuroscience Institute Cavalieri Ottolenghi (NICO), approved by the Ethical Committee of the University of Torino, and authorized by the Italian Ministry of Health (authorization number: 168/2020-PR). The experiments were performed in accordance with the European Community Parliament and Council Directives of 24 November 1986 (86/609/EEC) and 22 September 2010 (2010/63/EU). Mice were housed with a 12 h light/dark cycle and free access to food/water. Adequate measures were taken to minimize pain and discomfort. Female C57BL/6j mice used for all the experimental procedures were purchased from Envigo RMS srl (Udine, Italy).

Antibody Plasma Concentration Kinetic

RDO24_mIgG2a, RDO24_mFab2, or RDO24_mFab were intravenously injected in C57BL/6j mice (n=3) at the following concentration 10, 7 and 3.5 mg/kg, respectively. Plasma was collected after 1, 6, 24, 72 and 120 hours for each mouse and diluted 1:800 in PBS for antibodies quantification. 96well EIA/RIA plates (Costar) were coated at 4°C overnight with 100ng/well human MET ECD-Fc (R&D). Following three washes with PBS with 0.05% Tween 20, whole RDO24 mAb and fragments. Images of wound at the start moment and after the treatment were taken with DMRI Leica inverted microscope. Migration was quantified by evaluating the area of wound at time zero (A0) and at time after the treatment (Ay = 24 h). Normalization and quantification on the basis of three independent experiments were obtained by the formula (A0–Ay)/A0.

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Histological Evaluation

EAE induction and Clinical Evaluation

As reported in Montarolo et al., 2014, 2015, 2021 (50–52), to induce EAE, 6–8 week-old-female C57BL/6 mice were immunized by subcutaneous injection under the rostral part of the flanks and at the base of the tail with 300 µl of 2 µg/mouse of myelin oligodendrocyte glycoprotein (MOG35–55; Espikem, Florence, Italy) in incomplete Freund’s adjuvant (IFA; Sigma-Aldrich, Milan, Italy) containing 8 mg/mL Mycobacterium tuberculosis (strain H37Rv; Difco Laboratories Inc., Franklin Lakes, NJ, USA). Mice were treated with two intravenous injections of 500 ng of Pertussis toxin (Duotech, Milan, Italy) on the immunization day and 48 h later. Clinical score (0 = healthy; 1 = limp tail; 2 = ataxia and/or paresis of hind limbs; 3 = paralysis of hind limbs and/or paresis of forelimbs; 4 = tetraplegia; 5 = moribund or dead) was recorded daily by an investigator blind to group identity. The percentage of disease-free mice was calculated evaluating the day post immunization (dpi) when the first clinical manifestations appeared (score>0). Cumulative score was calculated as the sum of the daily score during experiment. To obtain the final concentration (10 mg/kg) RDO24 was dissolved in saline solution (0.9% NaCl). Control animals received the vehicle (0.9% NaCl, vehicle). The RDO24 or vehicle EAE mice were intravenously injected at 6, 8, and 10 days post immunization (dpi). The experiment with 5 vehicle- and 5 RDO24-treated animals was twice repeated.

Histological Evaluation

EAE mice were deeply anesthetized (zotetil 100 mg/kg, xylazine 5 mg/kg) and trans-cardially perfused with 4% paraformaldehyde in 0.12 M phosphate buffer, pH 7.2–7.4. The spinal cords were removed and immersed in the same fixative at 4°C for 24 h and then cryo-protected in 30% sucrose in 0.12 M phosphate buffer. Spinal cord were frozen and serially cut by a cryostat in 30 µm-thick coronal sections collected in PBS. For immunofluorescence, sections were stained to detect the expression of MOG (1:200, Proteintech, Rosemont, IL, USA) antigen. Incubation with primary antibody was made overnight at 4°C in PBS with 0.5% Triton-X 100. The sections were then exposed for 2 h at room temperature with secondary Alexa Fluor 555 (Molecular Probes Inc, Eugene, regon) -conjugated antibody. 4,6-diamidino-2-phenylindole (DAPI, Fluka, Saint Louis, USA) was used to counterstain cell nuclei. After processing, sections were mounted on microscope slides with Tris-glycerol supplemented with 10% Mowiol (Calbiochem, LaJolla, CA). Histological specimens were examined using ZEISS Axioscan 7 Microscope Slide Scanner (Weltzar, Germany). Adobe Photoshop 6.0 (Adobe Systems, San Jose, CA) was used to assemble the final plates. Quantitative evaluations were performed on images followed by ImageJ (Research Service Branch, National Institutes of Health, Bethesda, MD; available at http://rsb.info.nih.gov/ij/) analyses. The expression level of MOG staining as positive fractioned area (i.e. the percentage of positive pixels throughout the entire area)
was quantified to analyse the demyelinated areas. Inflammatory infiltrates were assessed using DAPI staining.

**Statistical Analysis**

All values are expressed as the mean ± standard deviation of, at least, 3 independent experiments. T-test was used to statistically compare two groups. For data representation in graph we performed data transformation expressing the values as ‘fold mean control’. For animal studies data were represented as mean ± standard error (SE) or median. Normality of distribution was assessed by the Shapiro-Wilk test. Two-way ANOVA followed by the Bonferroni’s multiple comparison post-hoc test was used to compare the clinical score during days. Mann–Whitney U test was used to compare continuous data between groups. Chi-square test was used to compare the onset of EAE. In all instances, the threshold P value deemed to constitute statistical significance was <0.05. The data analysis and the graph design were done using GraphPad Prism software.

**RESULTS**

**Engineering of the RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab**

We engineered the full-length form of mouse DO24 mAb and its derived recombinant fragments (Fab2 and Fab) (Figure 1A). The mAb cDNA was cloned from total RNA of the corresponding hybridoma using RT-PCR (reverse transcriptase-polymerase chain reaction) based on a degenerated oligonucleotide strategy. Then, using the mAb heavy and light chains as templates, the regions corresponding to the variable (VH and VL) domains, respectively, were PCR-amplified. Synthetic codon-optimized gene sequences for VH and VL chains were sub-cloned into relevant pcDNA3.1 vector expressing the murine constant IgG2a heavy chain (CH1, CH2, CH3) or the murine constant k light chain (CL), respectively. The recombinant Fab2 mAb was built by joining the VH-CH1 with the hinge sequence present on the vector, containing the cysteine residues required for dimerization. The recombinant Fab was obtained by expressing VH-CH1, VL-CL sequences. Moreover, the heavy chains of Fab and Fab2 fragments were tagged with a histidine tail for purification purposes. Figure 1B shows the drawing of the bivalent and monovalent structures of the molecules. The Figure 1C shows that purified recombinant proteins, analyzed under non-reducing conditions, formed covalently linked complexes of 150 kDa (full-length RDO24_IgG2a), 100 kDa (RDO24_mFab2), and 50 kDa (RDO24_mFab). When analyzed under reducing conditions, the full-length antibody showed two bands of 50 and 25 kDa corresponding to the heavy and light chains, respectively. The mFab2 and mFab showed the expected molecular weights of 25 kDa for the shorter heavy chains as well as the light chains.

![Figure 1](https://example.com/figure1.png)

**FIGURE 1** | Schematic representation (A), hypothesized structure (B), and western blot analysis (C) of RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab. NR, not-reduced; R, reduced.
RDO24 Antibody and Fragments Bind to the ECD of Human MET With High Affinity

The binding kinetics of RDO24 molecules to recombinant human MET extracellular domain-fragment crystallizable region (ECD-Fc) were determined by means of Surface Plasmon Resonance (SPR) using BiacoreT100. The specificity of the interaction between RDO24 and hMET ECD-Fc was evaluated by single cycle kinetic analysis. As described in the “Material and Methods” section, the human MET ECD-Fc was directly immobilized on a CM5 sensor chip by amine coupling. Then, increasing concentrations of RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab were passed onto the sensor chip, and the kinetic of their interaction with MET was measured with respect to the antigen association ($k_{on}$) and dissociation rate ($k_{off}$), from which we calculated the dissociation constant ($k_D$) of each construct toward the MET ECD-Fc target. The resulting SPR sensorgrams revealed a high affinity interaction between human MET ECD-Fc and RDO24_mIgG2a or RDO24_mFab or RDO24_mFab2, with $k_D$ value of 8.34 nM, or 8.85 nM or 7.08 nM, respectively (Figure 2). Moreover, the kinetic profile of the tested fragments revealed a low dissociation rate supporting a highly stable binding with the MET receptor.

RDO24 Cross-Reacts With Human, Rat and Mouse c-MET

By using flow cytometry experimental approach, we demonstrated that RDO24 antibody binds the MET receptor of human, rat and mouse origin (Figure 3). RDO24 mAb recognized the surface MET protein expressed in different types of human cells: GTL16 gastric carcinoma, A549 lung adenocarcinoma, HEK293T human embryonic kidney cells, and HUVEC human umbilical vein endothelial cells (Figure 3). MAbs interacted also with the MET receptor of rat cardiomyoblast H9c2 and mouse rhabdomyosarcoma cell lines (Figure 3).

Mapping of the RDO24 Epitope on the PSI Domain of MET Receptor

The extracellular region of the MET receptor consists of a semaphorin (SEMA) domain, a plexin semaphorin-integrin (PSI) domain (similar in structure to the plexins, semaphorins and integrins), and four immunoglobulin-like regions found in plexins and transcription factor (IPT) (MET WT in Figure 4A). The SEMA domain consists of a seven-blade β-propeller fold: the blades 1-4 comprise the MET α-chain while the blades 5-7 are part of the β-chain. The full-length single-chain MET precursor, in fact, is cleaved into two disulfide-linked chains of 50 kDa α- and 145 kDa β- subunits in the Golgi. The DO24 antibody was previously shown to bind the extracellular portion of MET β-chain (45), but its precise binding site was not determined. In order to precisely map the MET epitope recognized by RDO24 mAb, we used MET WT and different engineered mutant molecules constituted by deleted portions of the extracellular domains fused to transmembrane and intracellular domains of the receptor: MET ΔPSI-ΙPT, MET ΔΙPT, MET ΔSEMA, MET ΔSEMA-PSI (Figure 4A). All proteins contain the endogenous leader sequence at the N-terminus. The constructs were transfected in TOV-112D cells, which do not express MET. As shown in Figure 4B, RDO24 mAb was able to immunoprecipitate full-length MET WT, resolved in the β-chain (145kDa) and α-chain (50kDa) under reducing conditions. Among the deleted proteins, RDO24

![FIGURE 2](image-url) | Binding kinetics of RDO24_mIgG2a, RDO24_mFab2 and RDO24_mFab. Human MET ECD-Fc was immobilized on the sensor chip, and binding of increasing concentrations of engineered antibodies was determined by surface plasmon resonance.
immunoprecipitated the MET ΔIPT (80 kDa β-chain and 50 kDa α-chain) and MET ΔSEMA (110 kDa β-chain), but not the MET ΔPSI-IPT or MET ΔSEMA-PSI mutant proteins. The MET ΔPSI-IPT migrated as a 120 kDa single-chain protein, suggesting that it was not cleaved into the αβ heterodimer. These data indicate that RDO24 likely recognizes an epitope located in the PSI domain of the MET receptor.

RDO24 Binds the ECD of MET and Not That of Closely Related RON and SEMA4D

The PSI domain is structurally conserved in MET family members such as recepteur d’origine nantais (RON) (53) and Semaphorin4D (SEMA4D) (54). To evaluate whether RDO24 specifically binds to MET, we performed ELISA binding assays with MET, RON and SEMA4D ECD. We showed that RDO24 binds the ECD of MET (Figure 5A) and not that of closely related RON (Figure 5B) and SEMA4D (Figure 5C), indicating the selectivity of RDO24_mIgG2a against MET protein.

RDO24 Docks Onto the PSI Domain of MET in Computational Modeling Analysis

A computational modeling was performed in order to dock RDO24 Fv fragments onto the MET structure available in the Protein Data Bank (PDB) online archive with the code 1SHY (55). We generated a three-dimensional structural model of Fv RDO24 using the Rosetta-based computational homology modeling technique (Figure 6A) (48). Next, we used the protein-protein docking program ZDOCK (49) to dock the Fv RDO24 on top of the 1SHY structure of MET. HGF binds the SEMA domain (light pink) of MET (Figure 6B). Superposition of RDO24 Fv fragments with MET showed that the CDR-H1, CDR-H3 and CDR-L2 strongly interact with PSI domain (yellow) of MET (Figures 6B, C).

RDO24_mIgG2a and RDO24_mFab2 Are Endowed With MET Agonistic Activity

The DO24 mAb was previously shown to promote MET phosphorylation and therefore acts as an agonist of the receptor (46). This is likely due to the bivalent nature of the antibody, which enables the MET receptor dimerization, transphosphorylation, and activation of the downstream signaling. To verify this hypothesis the MET activation potency of bivalent (whole and mFab2) and the monovalent (mFab) molecules was assessed. A549 cells were treated with either molecule, lysed and analyzed in western blot (Figure 7). HGF was used as positive control. The RDO24_IgG2a and mFab2, but not the mFab, induced phosphorylation of MET at Y1234/1235 in a dose-dependent manner (Figures 7A, B). Furthermore, RDO24_IgG2a and mFab2, but not the mFab, stimulated phosphorylation of ERK (extracellular signal regulated kinase) and AKT, the two main signaling pathways downstream MET, to a similar extent to that induced by HGF (Figure 7B). The downstream ERK effector, CREB (cAMP response element binding protein) transcription factor, was also phosphorylated after treatment with bivalent antibodies (Figures 7B). Time course experiments revealed that the bivalent RDO24 molecules showed MET phosphorylation responses that were comparable in time to the ones elicited by DO24 monoclonal antibody, from which they were derived, or by the natural HGF ligand (Figures 7C, D). These data indicate that the RDO24_IgG2a and Fab2 fragment own agonistic properties for MET leading to stimulation of receptor and its downstream signaling pathways.

FIGURE 3 | RDO24_mIgG2a cross-reacts with human, rat, and mouse MET receptor. MET surface protein levels were measured by flow cytometry in human GTL16, A549, HEK293T, and HUVEC, in rat H9c2, and in mouse rhabdomyosarcoma cells by using RDO24_mIgG2a labelled by PE-Cy7.

![Flow Cytometry Image](image-url)
RDO24_mIgG2a and RDO24_mFab2 Mimic the HGF-Mediated Biological Effects

One of the unique biological effects induced by HGF in epithelial cells is cell-cell dissociation. In fact, HGF (also known as scatter factor) was originally discovered as a fibroblast-derived factor able to induce cell scattering (2). As shown in Figure 8A, incubation of MDCK epithelial cells with HGF induced a morphological change from epithelial colonies into scattered individual cells. Importantly, both the entire and the Fab2 mAbs, but not the Fab fragment, exerted a scattering effect on MDCK cells, comparable to that of HGF (Figure 8A). For cell motility assay, HPAF-II cells were treated with MET agonists and analyzed by the XCelligence Systems for 24 h. Activation of MET receptor by HGF induced strong cell migration as compared to untreated cells (Figure 8B). Both full-size mAb and Fab2 produced a significant biological effect over control, though at lower extent as compared to HGF (Figure 8B). In a ‘wound-healing’ assay, treatment with either bivalent molecule induced rat H9c2 cardiomyoblast cells to migrate and to cover the wounded area to the similar extent as HGF (Figure 8C). Overall, these data suggest that both entire and Fab2 mAbs mimic the migratory effects of HGF.

RDO24 Delays the Onset of Clinical Symptoms in EAE Mouse Model In Vivo

To evaluate the stability of the new molecules in blood circulation in vivo, a single dose of RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab was injected intravenously and the plasma concentrations were determined by ELISA assay. As shown in Figure 9A, the full-size antibody was very stable whereas the Fab2 and Fab molecules were lost 6 hours after the injection. These data suggested that the full-size antibody was a more suitable molecule for in vivo treatments. To evaluate the effect of RDO24 on the onset of MOG35-55-induced EAE, immunized mice received the compound intravenously three times once every two days beginning from the 6th dpi, when phenotypic EAE signs are not yet evident, but the immunization process has already occurred. RDO24 treatment was able to significantly delay the disease onset as reported by analysis of clinical score during days (Figure 9B) and percent of disease-free mice (Figure 9C). The cumulative score of RDO24-treated EAE mice was also significantly reduced in comparison to vehicle (Figure 9D). Mice were sacrificed in the 21th dpi, and the spinal cords were stained with MOG antibody for the assessment of demyelination in the white matter (Figure 9E). No significant
DISCUSSION

MET is a tyrosine kinase receptor that is physiologically activated by its unique HGF ligand. Activation of MET promotes cell proliferation and survival, is involved in tissue protection and repair, and represents an interesting target for regenerative medicine. Unfortunately, despite its therapeutic potential, HGF is a poor drug candidate. Exogenous HGF administered by intravenous injection is rapidly cleared by the liver in vivo (56, 57). Moreover, HGF is sequestered by the low affinity–high avidity sites widespread among the extracellular matrix proteoglycans (58–60). Additionally, as the endogenous HGF is post-translationally modified and correctly folded after its biosynthesis, the industrial production of bioactive HGF is hampered by the difficulty of producing a fully active molecule. Hence, the clinical use of HGF is limited and its substitution with molecules mimicking HGF, such as agonist monoclonal antibodies, is justified. Monoclonal antibodies are useful human therapeutics because of their specificity, affinity, and structure stability. Furthermore, they can easily be engineered in vitro using synthetic genes (61).

HGF is produced and secreted as an inactive single-chain pro-HGF form. Proteolytic cleavage is required to obtain the disulfide-bond linked two-chains HGF, which is the active form able to activate MET. MET can also be activated by bypassing the proteolytic processing of single chain pro HGF through an allosteric mechanism involving peptides. In particular, the so-called peptide V8 was found to bind the single-chain pro-HGF resulting in a similar conformation of two-chain HGF observed in the activation cleavage pathway, representing a new approach for MET signaling activation (62).

Another strategy to mimic HGF activity is the generation of artificial MET bivalent macrocyclic peptides, covalently bound by linkers such as Polyethylene Glycol (PEG) or carbon chains. Bivalent macrocyclic peptides can dimerize and activate MET depending on the linker length and show a full agonism towards MET, thus mimicking HGF activation in a similar extent (63). A further approach is the engineering of HGF fragments. HGF is composed of an N-terminal hairpin domain, four Kringle domains, and a C-terminal serine protease homology domain (64). Even if the N-terminal and first kringle fragment (NK1) occurs as a natural variant able to activate MET receptor, it is limited by its low stability and weak agonist activity. NK1 mutants were thus engineered to achieve a better stability and higher agonist potency and disulfide linked NK1 homo dimers showed a similar agonistic activity to that of full-length HGF (65). Furthermore, a modern approach is the targeting of MET receptor with antibodies. Herein, we describe the engineering and characterization of a monoclonal murine antibody acting as MET receptor agonist that might be useful for regenerative medicine.

For this purpose, we generated the new recombinant RDO24_mIgG2a and mFab2 fragment by genetic engineering. The RDO24_mIgG2a and mFab2 were created to produce bivalent molecules with a unique variable portion (VH-VL) interacting with the cognate MET antigen. These molecules contain two antigen binding portions (VLCL and VHCH1) linked together by a disulphide bond. The presence of two identical monovalent binding sites may give to the antibody molecules the agonistic property. For this reason, we tested whether recombinant full-length and Fab2 molecule mimic the biological effects exerted by the natural HGF ligand in molecular, cellular, and biological assays. There are many advantages of using Fab2 fragment antibodies instead of whole IgG antibodies. Fab2 fragment antibodies do not have Fc portions, thus they eliminate non-specific binding between Fc portions of antibodies and Fc receptors on cells (such as macrophages, dendritic cells, neutrophils, NK cells and B cells). Indeed, for a stimulating antibody to be used in regenerative medicine it is an advantage in avoiding the Fc-mediated functions such as antibody-dependent cell-mediated cytotoxicity or complement-dependent cytotoxicity.
Another advantage is that Fab2 fragments penetrate tissues more efficiently than whole IgG antibodies due to their smaller size. Last, but not least the smaller size of fragments may permit cheaper, faster production in microbial systems. On the other hand, the smaller size of Fab2 fragments may give rise to shorter half-life and increase renal clearance, representing a limitation for the development in therapy.

HGF binding upon MET leads to its dimerization, which is mandatory for its activation. Activation occurs by trans-phosphorylation of the two catalytic tyrosines Y1234-Y1235 belonging to the tyrosine kinase domain. Two docking tyrosines (Y1349-Y1356) of the C-terminal domain are subsequently phosphorylated and signal proteins, such as Gab1 (GRB2-associated-binding protein 1), Src, PI3K (phosphoinositide-3-kinase), GRB2 (Growth factor receptor-bound protein 2), PLCγ (phospholipase c gamma) and STAT3 (signal transducer and activator of transcription 3), containing a binding site for Src-homology-2 (SH2) or a phosphorylation domain, are recruited (44). These events lead to the activation of MET effectors, such as AKT and RAS-MAPK (mitogen-activated protein kinase) pathway, which are required for MET biological functions. We found that stimulation of A549 cells with bivalent whole antibody and Fab2 fragment induced MET phosphorylation at Y1234/1235 and, consequently, enhanced the levels of P-AKT and P-ERK, which are the main signaling downstream of MET. Furthermore, the transcription factor CREB, an effector of ERK, was also phosphorylated. Stimulation with monovalent Fab did not exert the same results. Thus, a bivalency-driven mechanism of action confers agonistic properties to RDO24_mIgG2a and mFab2 for MET activation and induction of its downstream signaling pathways. In line with this, the bivalent but not the monovalent mAbs evoke MET triggered biological effects, including cell scattering, migration, and wound healing. Our mAb cross reacts with mouse and rat MET. This represents an important requirement for pre-clinical models of regenerative medicine, requiring the antibody to be employed on rodent tissues and cells.

To test the potentially promising perspective of RDO24 agonist molecules to reach the preclinical phase, we performed in vivo analysis in mice. Unfortunately, the Fab2 fragment showed a short half-life in blood circulation. Thus, we considered the more stable full-size antibody for preclinical studies. We used the EAE mouse model of MS, a disease characterized by leucocytes infiltration and accumulation in the central nervous system and local destruction of myelin and neurons. Previous work showed that HGF exerts anti-inflammatory and immunosuppressive functions, showing efficacy in the mitigation of EAE (38, 66). Our results in vivo suggest that RDO24 is a functional HGF mimetic and may be beneficial in inflammatory diseases. However, there is an important caveat for pharmacological intervention at the level of MET activation to protect from inflammation and promote tissue regeneration. In fact, MET is involved in cancer progression, and some tumors are driven by MET alterations, such as amplification and overexpression (22). Further studies are necessary to ultimately elucidate the window of therapeutic benefit, without inducing adverse effects.

The MET extracellular domain is composed by three types of structural domains: SEMA, PSI, and IPT (1–4). The SEMA domain is shown in light pink, the PSI domain in yellow and HGF in green (B); enlarged view in (C).
Domain is structured as a seven-bladed propeller, whose blades 1-4 comprise the MET α-chain while blades 5-7 belong to the β-chain. HGF consists of two α and β subunits linked by a disulphide bond, as mentioned in the introduction. The HGF α-chain contains the high-affinity MET binding site (8) and interacts with SEMA domain (67) and also with IPT3,4 regions (13). The cleaved β-chain of HGF was crystallized together with MET SEMA-PSI domains (55) and interacts with low affinity with the bottom face of blades 2 and 3 of SEMA domain. While the HGF α-chain is sufficient for MET binding, the cleaved β-chain is necessary for MET activation (55, 68). The MET PSI domain does not interact with HGF. Although its function is not known, circumstantial evidence suggests that it may work as an hinge and likely orients the adjacent domains SEMA and IPT for proper ligand binding (69, 70).

We previously showed that DO24 binds to the extracellular portion of MET β-chain and does not compete with HGF binding (46). A SPR analysis revealed a high affinity and stable interaction of our recombinant full-length mAb and its fragments, with human MET ECD-Fc, with an equilibrium kD in the low nanomolar range, and a kinetic profile typical of highly stable protein-protein interactions. Furthermore, deletion mapping studies showed that the binding of RDO24 to the MET ECD was abolished by deletion of the PSI domain. The involvement of PSI domain in its binding to MET was confirmed by computational modeling. The PSI domain is structurally conserved in MET family members such as RON (53) and SEMA4D (54). Our mAb recognizes the PSI domain of the MET ECD but not the same domain of RON and SEMA4D receptors, belonging to MET family, thus suggesting a selectivity towards MET. Different monoclonal antibodies have been raised to the extracellular domain of MET (see Prat et al., 2014 for a review) (71). The bivalent antibodies directed to the SEMA domain usually behave as agonists, as the SEMA contains the low and high affinity binding sites for HGF (11) and is critical for receptor dimerization and activation (12). The bivalent agonist antibody (5D5) against the SEMA domain competes with HGFβ binding site and was converted to a monovalent

![FIGURE 7](image-url)
one-armed human IgG1 format to become antagonist (72). The DN30 antibody, endowed with partial agonist activity, proved to be a potent antagonist when engineered to a monovalent Fab (73). The MET epitope recognized by DN30 is within the IPT4 domain (74), a region included in one of the identified HGF binding sites (domains IPT3 IPT4) (13). Notably, the “antagonist” mechanism of DN30 is associated to induction of proteolytic cleavage of MET ECD (receptor “shedding”) followed by proteasome-mediated receptor degradation (73). In both cases, the conversion of a bivalent antibody into a monovalent form has been used to minimize the agonist activity. The role of PSI domain in the regulation of MET is still unknown. Recently, an antagonistic antibody specific to the PSI domain of MET, ICR201, has been reported (75). ICR201 is a human IgG1 bivalent antibody, which induces rapid depletion of MET protein via the lysosomal degradation pathway and inhibits tumor growth in vitro and in vivo. Our bivalent anti-PSI antibody is a mouse IgG2a and is endowed with potent agonist activity. It is plausible that the IgG architecture contributes to the antagonist/agonist activity of

![FIGURE 8](image)
the antibodies (76). Future work will show whether constructing a human bivalent and/or monovalent one-armed IgG1 RDO24 antibody may be effective to create an inhibitory anti-MET antibody for cancer therapy.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/supplementary material. Further inquiries can be directed to the corresponding author.

**AUTHOR CONTRIBUTIONS**

CD contributed to figures realization, performed the ELISA assays and the docking study, and wrote the manuscript with TC and SG. SG and AV performed the experiments and realized the figures. EV cloned DO24 RNA hybridoma. CB and FZ realized MET mutant constructs and their transduction in TOV112D cells. FM performed treatments and analysis on EAE mice. EC performed the flow cytometry assays. RM and DMF performed SPR-based analysis. AB and PMC revised the manuscript and TC engineered the antibodies and wrote and revised the manuscript. All authors contributed to the article and approved the submitted version.

**FUNDING**

This work was supported by Fondazione Italiana Sclerosi Multipla (FISM), AIRC Sperimille Program N.21719, AIRC IG N. 23820 (grant number 2017/R/9) and by Italian Ministry of Health “Ricerca Corrente 2021”.

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**FIGURE 9** | RDO24 administration delays the onset of EAE. RDO24_mIgG2a, RDO24_mFab2, and RDO24_mFab plasma concentration kinetics were evaluated by ELISA after intravenous (I.V.) delivery in mice (A). The clinical course of EAE in RDO24 (n=10) and vehicle (n=10) –treated EAE mice was compared as mean clinical score (B), percentage of disease free mice (C), and cumulative (D) score. Dashed line indicates the days of treatment (6, 8, and 10 days post immunization). (E) Representative immunofluorescence images of coronal sections of spinal cord of vehicle (n=5) and RDO24 (n=5) -treated EAE mice stained with anti-MOG antibody (red). DAPI (blue) counterstains cell nuclei. Demyelination and perivascular inflammatory infiltrates were measured using respectively MOG fractioned area (F), and DAPI+ perivascular inflammatory infiltrates density (G) in vehicle- and RDO24-treated EAE mice. White arrows indicate the perivascular inflammatory infiltrates. Calibration bars, 100µm. Two Way ANOVA, Bonferroni post-test (B); Chi square test (C); Mann-Whitney U test (D, F, G). **Pvalue < 0.01 significant vs Vehicle; *Pvalue < 0.05 significant vs Vehicle.
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