BYON4228 is a pan-allelic antagonistic SIRPa antibody that potentiates destruction of antibody-opsonized tumor cells and lacks binding to SIRPg on T cells

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ABSTRACT

Background Preclinical studies have firmly established the CD47-signal-regulatory protein (SIRPa) axis as a myeloid immune checkpoint in cancer, and this is corroborated by available evidence from the first clinical studies with CD47 blockers. However, CD47 is ubiquitously expressed and mediates functional interactions with other ligands as well, and therefore targeting of the primarily myeloid cell-restricted inhibitory immunoreceptor SIRPa may represent a better strategy.

Method We generated BYON4228, a novel SIRPa-directed antibody. An extensive preclinical characterization was performed, including direct comparisons to previously reported anti-SIRPa antibodies.

Results BYON4228 is an antibody directed against SIRPa that recognizes both allelic variants of SIRPa in the human population, thereby maximizing its potential clinical applicability. Notably, BYON4228 does not recognize the closely related T-cell expressed SIRPg that mediates interactions with CD47 as well, which are known to be instrumental in T-cell extravasation and activation. BYON4228 binds to the N-terminal Ig-like domain of SIRPa and its epitope largely overlaps with the CD47-binding site. BYON4228 blocks binding of CD47 to SIRPa and inhibits signaling through the CD47-SIRPa axis. Functional studies show that BYON4228 potentiates macrophage-mediated and neutrophil-mediated killing of hematologic and solid cancer cells in vitro in the presence of a variety of tumor-targeting antibodies, including trastuzumab, rituximab, daratumumab and cetuximab. The silenced Fc region of BYON4228 precludes immune cell-mediated elimination of SIRPa-positive myeloid cells, implying anticipated preservation of myeloid immune effector cells in patients. The unique profile of BYON4228 clearly distinguishes it from previously reported antibodies representative of agents in clinical development, which either lack recognition of one of the two SIRPa polymorphic variants (HEFLB), or cross-react with SIRPg and inhibit CD47-SIRPg interactions (SIRPAB-K322A, 1H9), and/or have functional Fc regions thereby displaying myeloid cell depletion activity (SIRPAB-K322A). In vivo, BYON4228 increases the antitumor activity of rituximab in a B-cell Raji xenograft model in human SIRPa+ transgenic mice. Finally, BYON4228 shows a favorable safety profile in cynomolgus monkeys.

Conclusions Collectively, this defines BYON4228 as a preclinically highly differentiating pan-allelic SIRPa...
antibody without T-cell SIRPγ recognition that promotes the destruction of antibody-opsonized cancer cells. Clinical studies are planned to start in 2023.

**BACKGROUND**

Antibodies against tumor-associated antigens (anti-TAAs) play a prominent role in the treatment of a broad range of solid and hematologic cancers.1 Some of the most commonly used examples of anti-TAAs include trastuzumab directed against Her2 (over)expressed on breast cancer, cetuximab directed against the epidermal growth factor receptor (EGFR) on various carcinoma cells, rituximab directed against CD20 expressed on malignant B cells, and daratumumab directed against CD38 on multiple myeloma cells. Generally, anti-TAAs act by a combination of target-related direct and immune-mediated mechanisms. The immune-mediated mechanisms include complement-dependent cytotoxicity (CDC), effector cell-mediated antibody-dependent cellular cytotoxicity (ADCC) performed by natural killer (NK) cells and granulocytes, and antibody-dependent cellular phagocytosis (ADCP) exerted by macrophages.2-4 Furthermore, anti-TAAs can also effectively trigger adaptive T cell-mediated immunity by facilitating cross-presentation of tumor antigens to cytotoxic T lymphocytes.5 All these immune effector cell responses are triggered by the Fc regions of anti-TAAs that ligate activating Fc receptors and thereby initiate intracellular signaling and (direct or indirect) downstream tumor cell elimination. Despite this multitude of mechanisms, there remains a pertinent need to improve anti-TAAs clinical efficacy.

The CD47-signal-regulatory protein (SIRP)α axis has been firmly established as a myeloid immune checkpoint in preclinical and in early stage clinical studies.8-12 SIRPα is a typical inhibitory immunoreceptor primarily expressed on myeloid cells, including monocytes, macrophages, granulocytes and dendritic cell subsets.15-16 Two polymorphic variants are present in the human population, named SIRPαUT (also known as V1) and SIRPαS (also known as V2). SIRPα is the only inhibitory member of a multigene receptor family with the closest homologs in humans and other primates being SIRPβ1v1, SIRPβ1v2 and SIRPγ1. CD47, also known as integrin-associated protein, is the cellular ligand for SIRPα and SIRPγ, but not for the two SIRPβ1 receptors.18-22 CD47 is broadly expressed on most if not all cells in the body and is often found to be overexpressed on cancer cells.9,10 Cellular CD47 binding to SIRPα triggers inhibitory intracellular signaling via immunoreceptor tyrosine-based inhibitory motifs in the SIRPα cytoplasmic tail involving the recruitment and activation of the tyrosine phosphatases SH2 domain-containing protein-tyrosine phosphatases (SHP)-1 and/or SHP-2, which restricts myeloid effector function.13,15,16,23-26 Consequently, blockade of CD47-SIRPα signaling can promote macrophage-mediated and neutrophil-mediated tumor cell destruction in the presence of cancer-opsonizing antibodies, or other pro-phagocytic signals.9-11,27-30 In addition, there is accumulating evidence that disruption of the CD47-SIRPα signaling axis promotes adaptive anticancer immunity31,32 in combination with programmed cell death protein-1 (PD-1)/programmed death ligand-1 checkpoint inhibitors,33,34 or other approaches such as radiotherapy that can also trigger antitumor immunity.35

Around 35 therapeutics directed towards the CD47-SIRPα axis have entered clinical trials in recent years.36-38 The first clinical studies were performed with CD47-targeting agents, used as single agents or in combination with anti-TAAs or anti-PD-1, and have shown limited toxicity and promising initial efficacy.36,37,39-41 However, conceptually there appear to be several disadvantages of targeting CD47 per se. First, CD47 is not specific for tumor cells. In fact, CD47 is widely distributed, therefore forming a large ‘antigen sink’ requiring high doses of drug for saturation. Furthermore, many CD47-targeting agents have Fc tails with normal effector functions, including Fc receptor binding, and this may opsonize normal cells and cause toxicity, for instance, by promoting the immune-mediated destruction of such healthy cells. Indeed, anemia and thrombocytopenia are common side effects of CD47-targeting agents with functional Fc regions, often requiring red blood cell transfusion, even though a low-dose priming strategy has been adopted to mitigate this to some extent.36,39 Finally, CD47 does not only bind SIRPα but it also mediates functional interactions with integrins,42,43 vascular endothelial growth factor-144, thrombospondin-145 and SIRPγ,46,47 these may be affected by CD47-targeting agents too. The most notable of these other CD47 ligands is SIRPγ, a close homolog of SIRPα present only in primates, which is expressed on T cells and activated NK cells.45 CD47-SIRPγ interactions are pivotal for T-cell extravasation and activation and their disruption might therefore curtail durable antitumor immunity.48,46,47,48 Thus, targeting of CD47 may not be the most optimal way to selectively antagonize CD47-SIRPα interactions and therefore therapeutic targeting of the myeloid cell-restricted SIRPα may represent a better strategy.

Currently, four therapeutic SIRPα-targeting antibodies have entered clinical development in cancer indications: BI 765063 (NCT03990293, NCT04633142, NCT05249426, NCT05461299), BI 770371 (NCT05327946; no details on the antibody nor its properties have been disclosed yet), BMS-986351 (NCT03783403, NCT05168202), and GS-0189 (NCT04502706) (also see table 1). However, most of the anti-SIRPα antibodies HEFLB, 1H9, and SIRPAB-11, which in all probability are representative for three of the clinical stage SIRPα antibodies (see table 1), have been reported to lack binding to both polymorphic SIRPα variants that are present in the human population, or they also recognize the related SIRPγ.49

Here we report the preclinical characterization of BYON4228 and a direct in vitro comparison with the three anti-SIRPα antibodies HEFLB, 1H9 and SIRPAB-11.
BYON4228 is a pan-allelic SIRPα antibody that lacks binding to SIRPγ present on T cells. BYON4228 binds to a region overlapping with the CD47 binding site on SIRPα, thereby preventing binding of CD47 to SIRPα and blocking inhibitory signaling. BYON4228 potentiates killing and phagocytosis by human neutrophils and macrophages, respectively, of hematologic and solid tumor cells opsonized with various anti-TAAs. Silencing of the BYON4228 Fc tail prevents unwanted destruction of SIRPα-positive myeloid cells. In a human SIRPαζ+ transgenic mouse model, BYON4228 promotes rituximab-dependent elimination of Raji cells. Finally, intravenous single dose infusion of up to 100 mg/kg and repeated dose infusion of up to 30 mg/kg BYON4228 to cynomolgus monkeys is well tolerated. These data support further development of BYON4228 in clinical trials.

**METHODS**

Detailed methods can be found in the online supplement.

**BYON4228 antibody development and humanization**

BYON4228 was generated by immunization of rabbits with the extracellular domains (ECDs) of SIRPα allelic variants, sequencing PCR products from selected B cells and humanization.

**HEFLB, SIRPAB-11-K322A, 1H9 antibody sequences**

The sources of the antibody amino acid sequences of HEFLB, SIRPAB-11-K322A (referred as SIRPAB-11 in this paper) and 1H9 (the latter generated as a G1m17,1 allotype) (also see online supplemental materials) and their presumed equivalents in clinical trials are shown in **Table 1**.

**Anti-TAAs**

The following anti-TAAs were used: rituximab (MabThera, Roche), daratumumab (DARZALEX, Janssen Biologics), panitumumab (Vectibix, Amgen), cetuximab (Erbitux, Merck) and trastuzumab (Herceptin, Roche).

**Table 1** Properties of anti-SIRPα antibodies used for in vitro comparisons

<table>
<thead>
<tr>
<th>mAb</th>
<th>Source of sequences</th>
<th>Fc tail</th>
<th>Names of corresponding agent in clinical trials*</th>
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<tbody>
<tr>
<td>HEFLB</td>
<td>WO 2017/178653</td>
<td>IgG4-S228P/L445P</td>
<td>Bl-765063 OSE-172</td>
</tr>
<tr>
<td>SIRPAB-11-K322A</td>
<td>WO 2020/068752</td>
<td>IgG1-K322A</td>
<td>BMS-986351 Anzurtofart CC-95251</td>
</tr>
<tr>
<td>1H9</td>
<td>WO 2019/023347</td>
<td>IgG1-N297A</td>
<td>GS-0189 FSI-189</td>
</tr>
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</table>

*Based on data provided in Gauttier et al. WO 2017/178653 and WO 2019/175218,87 HEFLB most likely corresponds to Bl-765063/OSE-172; based on identity of amino acid sequences in the publication of the International Non-proprietary Name (WHO Drug Information Vol 36 No 2 2022, p319) and WO 2020/068752. SIRPAB-11-K322A corresponds to anzurtofart/BMS-986351/CC-95251; based on data provided in Liu et al. Corporate presentation of 2020, 1H9 most likely corresponds to GS-0189/FSI-189.

**Cellular binding to SIRP-expressing Expi Chinese hamster ovary-S cells and primary granulocytes**

Transiently transfected Expi Chinese hamster ovary-S (ExpiCHO-S) cells were first incubated for 30 min with primary antibodies and then with AF647-conjugated or R-phycocerythrin (PE)-conjugated anti-human IgG antibodies. Median fluorescent intensities were determined by flow cytometry (FACSymphony or FACSVerse, BD Bioscience) or relative fluorescence units were determined (EnVision, PerkinElmer).

**CD47-bead and mAb binding to U937 cells and primary cells**

Peripheral blood mononuclear cells (PBMCs) or U937 cells were stained with Alexa Fluor 647-labeled antibodies and then incubated with CD47-bead and mAb binding to U937 cells and primary cells. Median fluorescent intensities were determined by flow cytometry (FACSsymphony, FACSVerse, BD Bioscience).}

**ADCP and ADCC**

ADCP was performed using confocal microscopy (CD20-positive target cells) or live cell imaging (CD20-negative target cells). Monocytes were differentiated to macrophages with macrophage colony-stimulating factor for 7 days. Fluorescently labeled or pHrodo-labeled target cells and antibodies were added for 3–8 hours at 37°C in the presence of 100 µg/mL intra-venous immunoglobulins (IVIG). For confocal ADCP, the cells were counterstained with anti-CD19-PE and Hoechst 33342 to determine phagocytosis (ImageXpress, Molecular Devices). For live cell imaging based ADCP, images were taken every 30 or 60 min using a real-time fluorescence imager (Incucyte, Sartorius). For ADCC, human neutrophils were isolated, activated for 30 min with 10 ng/mL granulocyte-macrophage colony-stimulating factor (GM-CSF), incubated for 4 or 20 hours with 100 µCi Cr-51 labeled target cells (effector:target ratio of 50:1), and killing was determined by analyzing the supernatant in

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a scintillation counter (MicroBeta² Microplate Counter, PerkinElmer).

Raji xenograft in human SIRPα-wt-Severe combined immunodeficiency transgenic mice

Tumors were induced by subcutaneous (SC) injection of 2×10⁷ Raji cells into the flank of female human SIRPα-wt-severe combined immunodeficiency (scid) animals. BYON4228 was administered intraperitoneal (IP) three times a week for 4 weeks at a dose of 5 mg/kg. Rituximab was administered at 1 mg/kg IP.

Graphical representation, curve fitting and statistical analysis

Dose-response curves were fitted by non-linear regression with a variable slope (four parameters) in GraphPad Prism 9. EC₅₀ or IC₅₀ values were calculated in GraphPad Prism as the concentration that gives a response halfway between bottom and top of the curve. Statistical testing was performed in GraphPad Prism 9 as indicated in the figure legends. For functional assays (ADCC, ADCP), statistical analysis was performed on the ‘fold enhancement’ graphs. Since no values could be calculated for non-/poor-responsive donors, statistical tests were not performed on EC₅₀ and IC₅₀ summarizing graphs or figures 3B and 4C.

RESULTS

BYON4228 is a high affinity pan-allelic SIRPα antibody

We aimed to generate a blocking antibody against the inhibitory receptor SIRPα that binds to both the SIRPα-wt and SIRPα-polymorphic variants, but lacks binding to SIRPγ on human T cells. The blocking antibody should disrupt the binding of CD47 to SIRPα and inhibit CD47-induced SIRPα signaling. Rabbits were immunized with a mixture of proteins representing the ECDs of both SIRPα allelic variants and cynomolgus SIRPα. B cells from these rabbits were screened for antibody production levels, antigen binding and blocking to select B-cell clones producing antibodies that met our criteria. The variable domains of these B-cell clones were used to produce chimeric antibodies with hIgG1 backbone to allow further selection rounds. Ultimately, we selected BYON4228, a humanized antibody that met all our pre-set criteria, and compared it to three SIRPα antibodies that are considered to be representative of three clinical stage SIRPα antibodies (table 1). BYON4228 bound to both SIRPα-wt and SIRPα with high affinity (table 2, figure 1A,B). Clearly, this pan-allelic characteristic was not a common property of all other SIRPα-antibodies tested. In particular we found, in line with earlier publications,⁵¹ ⁵⁵ that the SIRPα antibody HEFLB did not display binding to SIRPα (figure 1A,B). All four anti-SIRPα monoclonal antibodies (mAbs) tested (ie, BYON4228, HEFLB, SIRPAB-11 and 1H9) showed binding to SIRPβ1v1 and SIRPβ1v2, although with variable EC₅₀ values (figure 1A,B). In line with its binding specificity, BYON4228 potently bound to primary SIRPα-expressing immune cells irrespective of their SIRPα-genotype (figure 1C,D). Overall, these results show that BYON4228 is a high-affinity pan-allelic SIRPα antibody that binds its native antigen on myeloid cells.

BYON4228 binds to a conserved epitope on the CD47-binding domain of SIRPα and blocks signaling

The extracellular region of SIRPα consists of three immunoglobulin superfamily (IgSF) domains and the first N-terminal V-set IgSF domain mediates binding to CD47.²¹ To identify the domain to which BYON4228 binds, we took advantage of the finding that BYON4228 lacks binding to SIRPγ (online supplemental figure S1A, table 3) and generated various chimeric SIRPα-SIRPγ domain swapped proteins. Binding of BYON4228 to these chimeric proteins could only be detected when domain 1 was derived from SIRPα, but not from SIRPγ. The origin of domain 2 and 3 was irrelevant for BYON4228 binding, showing that SIRPα domain 1 is both essential as well as sufficient for BYON4228 binding. Therefore, the BYON4228 epitope is present within this domain (online supplemental figure S1A). To define in more detail the epitope of BYON4228, we employed hydrogen deuterium exchange mass spectrometry technology. This method relies on the hydrogen to deuterium exchange, which occurs when molecules are incubated in heavy water. When a protein is complexed with an antibody, the deuterium uptake is hampered in the binding region, which can be measured using mass spectrometry. Using this method, BYON4228 binding signals were detected in the regions between residues 26–40, 50–62 and 95–103 (online supplemental figure S1B) of both SIRPα-wt and SIRPα. For visualization, the epitope was projected onto the N-terminal Ig-like CD47-binding domain of SIRPα-wt and SIRPα, previously resolved by X-ray crystallography.²¹ ⁵⁶ A large overlap with the CD47 binding site was noted (figure 2). The binding site covered almost exclusively non-polymorphic amino acid residues (online supplemental figure S1C), in line with the potent binding of BYON4228 to both SIRPα-wt and SIRPα. In conclusion, these results show that the BYON4228 epitope maps

![Graphical representation](image-url)
to the CD47 binding site on the membrane distal domain of SIRPα.

To test the ability of BYON4228 to disrupt the interaction between SIRPα and CD47, PBMCs were incubated with fluorescent beads coated with human CD47. The CD47-beads bound to SIRPα-expressing monocytes and BYON4228 was able to inhibit such binding in a dose-dependent manner, irrespective of the SIRPα-genotype (figure 3A,B). While all four tested anti-SIRPα antibodies showed binding to primary monocytes of all SIRPα-genotypes, HEFLB was unable to inhibit CD47 binding to monocytes of SIRPα1/α1 donors and inhibited CD47-bead binding to monocytes of SIRPαβ/α1 donors with limited efficacy (figure 3A,B). The binding

Table 3  Observed affinities of BYON4228 for binding to human SIRPγ

<table>
<thead>
<tr>
<th>Ligand for BYON4228</th>
<th>Observed affinities (K&lt;sub&gt;D&lt;/sub&gt;-obs)</th>
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<tbody>
<tr>
<td></td>
<td>Antigen on surface set-up</td>
</tr>
<tr>
<td>Human SIRPγ ECD</td>
<td>NB</td>
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*Dose-dependent very low binding responses were observed; no K<sub>D</sub> could be estimated.
ECD, extracellular domain; NB, no binding; SIRP, signal-regulatory protein.
BYON4228 and CD47 footprints projected on SIRPγ

Figure 2  BYON4228 binds to the CD47-binding domain of SIRPα. (A) The BYON4228 epitope was mapped using HDX-MS. Projection of the CD47 binding site21,56 (purple, left), BYON4228 HDX-MS mapped epitope (orange, middle) and overlap (brown, right) onto the N-terminal Ig-like CD47-binding domain of SIRPα and SIRPγ. HDX-MS, hydrogen deuterium exchange mass spectrometry; SIRP, signal-regulatory protein.

BYON4228 lacks binding to human T-cell expressed SIRPγ

The binding of BYON4228 and other antibodies to SIRPγ was then investigated. BYON4228 displayed no detectable binding to SIRPγ when transiently expressed on ExpiCHO-S cells and negligible/no binding to SIRPγ expressed as a soluble protein (figure 4A, table 3). In line with earlier reports,54 HEFLB also lacked binding to SIRPγ, whereas SIRPAB-11 showed potent binding to SIRPγ-expressing cells (figure 4A). We furthermore noted, in contrast to an earlier report,49 that 1H9 also displayed SIRPγ binding when expressed on ExpiCHO-S cells. In line with this, 1H9 also demonstrated binding to SIRPγ-expressing T cells, although with lower potency than SIRPAB-11 (figure 4B,C). Importantly, BYON4228 lacked binding to T cells altogether (figure 4B,C). CD47 is a known ligand for both SIRPα and SIRPγ55 and indeed, CD47-coated beads showed binding to T cells of most donors. The antibodies 1H9 and SIRPAB-11 blocked CD47-bead binding to T cells, while BYON4228 and HEFLB did not alter this binding (figure 4B,C). Overall, these results show that BYON4228 lacks binding to T-cell expressed SIRPγ and accordingly does not affect CD47 binding to SIRPγ on T cells.

BYON4228 promotes ADCC and ADCP of therapeutic mAbs in a pan-allelic fashion

We then continued to investigate the functional activity of BYON4228 with respect to tumor cell killing. We first studied the impact of BYON4228 on neutrophil-mediated ADCC induced by anti-TAAs. GM-CSF-activated neutrophils were able to kill SK-BR-3 breast cancer tumor cells opsonized with trastuzumab. Addition of BYON4228 potentiated this killing in a dose-dependent and pan-allelic fashion (figure 5A–C), with an average 2.5-fold to 3.0-fold enhancement (figure 5C) of trastuzumab-induced killing at saturating BYON4228 concentrations. ADCC enhancement by BYON4228 appeared most potent (ie, lower EC50 values) when neutrophils were of SIRPα/α1 origin (EC50 mean of 0.10 µg/mL vs 0.19 and 0.26 for SIRPα/α1 and SIRPα/α1 donors, respectively). However, at saturating levels of BYON4228, ADCC enhancement was not significantly different between different SIRPα genotyped donors (figure 5C).

As expected, the SIRPα/α1-specific antibody HEFLB was unable to potentiate trastuzumab-ADCC by SIRPα/α1 homozygous neutrophils. In addition, trastuzumab-ADCC enhancement of HEFLB by SIRPα/α1 neutrophils was limited (1.3-fold by SIRPα/α1 donors vs 3.0-fold by SIRPα/α1 donors). 1H9 potentiated ADCC by SIRPα/α1 neutrophils with limited efficacy (1.5-fold enhancement by SIRPα/α1 donors vs 3.3-fold enhancement by SIRPα/α1 donors), which was in line with its limited ability to antagonize CD47 binding to monocytes homozygous for SIRPα (figure 3A,B). To further evaluate the broadness of its therapeutic applicability, we also determined whether BYON4228 could enhance the ADCC activity of neutrophils towards other target cells, including cetuximab-opsonized and...
Figure 3  BYON4228 competitively blocks binding of CD47 to SIRPα. (A,B) anti-SIRP mAb binding and blockade of CD47 binding to primary monocytes. Indicated fluorescently labeled antibodies were incubated with primary peripheral blood mononuclear cells for 30 min, and then, fluorescently labeled CD47-coated beads were added. mAb binding and CD47 binding was measured on gated CD14-positive monocytes. (A) Top row depicts mAb binding and bottom row depicts CD47 binding to monocytes of representative donors with indicated SIRPα genotypes. (B) Overview of the mAb binding EC₅₀ values (geomeans are depicted), blocking of CD47 binding IC₅₀ values, and CD47 blocking windows (Δ MFIs) per antibody of all donors tested (seven SIRPα-BIT/BIT, five SIRPα-BIT/α₁, and six SIRPα-α₁ donors). No IC₅₀ values could be calculated from donors with incomplete curve saturation or no response as indicated in the graph. Donors that did not show a window for CD47 blocking were also indicated in the graph. (C) Graph depicts binding of CD47 coated fluorescent beads to U937 cells (left y-axis, open symbols), incubated in the presence of a dose-range of BYON4228-AF647 or AF647-labeled isotype control (right y-axis, closed symbols). Results display average binding MFIs ±SD measured in the FITC or AF647 channel of N=3 independent experiments. (D) SIRPα signaling measured using the PathHunter Jurkat SIRPα signaling reporter cell line (DiscoverX), after co-incubation with CD47-expressing or knock-out (KO) A431 cells, in presence of a concentration-range of indicated antibodies. Results are shown as mean ±SD of N=6 independent experiments. FITC, fluorescein isothiocyanate; mAb, monoclonal antibody; MFI, median fluorescent intensities; SIRP, signal-regulatory protein.
panitumumab-opsonized cancer cell lines (online supplemental figure S2). Both panitumumab (IgG2) and cetuximab (IgG1) alone were able to induce neutrophil ADCC towards the EGFR-expressing epidermoid tumor cell line A431 in a dose-dependent fashion (online supplemental figure S2A–C). BYON4228 enhanced the killing of both anti-EGFR antibodies (online supplemental figure S2E). We also tested panitumumab-induced and cetuximab-induced neutrophil ADCC of the colorectal cancer (CRC) cell line SW48 (online supplemental figure S3). Here, panitumumab consistently led to higher levels of neutrophil-induced killing when compared with cetuximab (4.5-fold vs 1.8-fold, respectively) (online supplemental figure S3C). Again, BYON4228 was able to enhance both panitumumab-induced and cetuximab-induced killing of the CRC cell line SW48 (1.7-fold vs 1.6-fold, respectively) (online supplemental figure S3E). We further tested isogenic SW48 cell lines containing frequently occurring constitutively activating mutations downstream the EGFR signaling pathway (Kirsten rat sarcoma virus (KRAS) G12D, KRAS G13D, rapidly accelerated fibrosarcoma B-type (BRAF) V600E). As expected,
Figure 5  BYON4228 enhanced ADCC and ADCP of therapeutic mAbs in a pan-allelic fashion. (A–C) Neutrophil-mediated ADCC measured using the Cr-S1 release assay after 4 hours incubation of target cells (SK-BR-3) and effector cells (primary granulocyte-macrophage colony-stimulating factor activated neutrophils) in the presence of a fixed dose of trastuzumab (10µg/mL) and a concentration range of indicated antibodies or respective isotype controls. Donors tested: Six SIRPαBIT/BIT, eight SIRPαBIT/α1, and six SIRPα1/α1 donors. (A) Results show concentration-dependent SIRPα-mAb induced % killing of SK-BR-3 cells by activated neutrophils of representative donors with indicated SIRPα genotypes. (B) SIRPα-mAb induced ADCC EC50 values of all donors. No EC50 values could be calculated from donors with incomplete curve saturation, no response or incorrect curve-fitting as indicated in the graph. (C) SIRPα-mAb induced ADCC fold enhancement of all donors. The means are depicted. The fold enhancement=(% killing at 10µg/mL trastuzumab + 4 µg/mL anti-SIRPα mAb)/(% killing at 10µg/mL trastuzumab). (D–F) Macrophage-mediated ADCP measured using confocal microscopy after 3 hours incubation of target cells (Raji) and effector cells (macrophages) in the presence of a fixed dose of rituximab (80 ng/mL) and a concentration range of indicated antibodies or respective isotype controls. Donors tested: Five SIRPαBIT/BIT, six SIRPαBIT/α1, and six SIRPα1/α1 donors. (D) Results show SIRPα-mAb induced Phagocytosis Index (PI) of macrophages from representative donors with indicated SIRPα genotypes. (E) SIRPα-mAb induced ADCP EC50 values of all donors. No EC50 values could be calculated from donors with incomplete curve saturation, no response or incorrect curve-fitting as indicated in the graph. (F) SIRPα-mAb induced ADCP fold enhancement of all donors. The means are depicted. The fold enhancement=(PI at 80 ng/mL rituximab + 10µg/mL anti-SIRPα mAb)/(PI at 80 ng/mL rituximab). For C and F, * p<0.05, ** p<0.01, ***p<0.001; p>0.05 is not indicated. P values were calculated by one-way analysis of variance (ANOVA) for each genotype (comparing the different mAbs) and for each mAb (comparing the different genotypes) followed by Tukey’s multiple comparisons test. ADCC, antibody-dependent cellular cytotoxicity; ADCP, antibody-dependent cellular phagocytosis; mAb, monoclonal antibody; SIRP, signal-regulatory protein; TAA, tumor-associated antigen.
the EGFR pathway mutant cell lines were killed by neutrophils after opsonization with panitumumab and this was further enhanced by BYON4228 (online supplemental figure S3F–H), although overall levels of killing were variable between the different cell lines, probably as a result of differences in EGFR expression levels (online supplemental table S1). The observation that BYON4228 was able to enhance the efficacy (i.e., maximum killing) of panitumumab in all tested cell lines (online supplemental figure S3F–H), implies that therapeutic efficacy of a combination of panitumumab and BYON4228 might be expected in patients that bear the indicated and other known growth-promoting mutations downstream the EGFR signaling pathway, an indication for which cetuximab and panitumumab are currently not registered/recommended.

The capacity of BYON4228 to enhance macrophage-mediated-ADCP induced by anti-TAAs was studied. CD20-expressing CellTrace Far Red labeled Raji cells (Burkitt’s lymphoma) were opsonized with rituximab and phagocytosis by monocyte-derived macrophages was assessed using confocal microscopy. To ensure phagocytosis, anti-CD19 was used as a counterstain and only CD19-negative Raji cells entirely engulfed by macrophages (i.e., truly phagocytosed) were counted. Rituximab-induced phagocytosis was increased by BYON4228 in a dose-dependent fashion (figure 5D–F). This synergistic effect was detected with all macrophages, irrespective of their SIRPα genotype. We noted that BYON4228 showed higher potency (i.e., lower EC_{50}) when macrophages were of SIRPα/α origin compared with SIRPα/̅α or SIRPα/̅α macrophages (EC_{50} of 0.05 vs 0.25 µg/mL, respectively) (figure 5E). However, the fold enhancement of the Phagocytosis Index of rituximab-induced phagocytosis by SIRPα/̅α donors at saturating 10 µg/mL was not significantly different when compared with SIRPα/̅α or SIRPα/̅α donors (figure 5F). The SIRPα/̅α-specific blocking antibody HEFLB was again not or only hardly capable of enhancing rituximab-induced phagocytosis by SIRPα-homozygous or heterozygous macrophages, respectively. Similarly, 1H9 showed limited efficacy and potency for phagocytosis enhancement by SIRPα/̅α and SIRPα/̅α macrophages. Remarkably similar results were found when Daudi cells (non-Hodgkin’s lymphoma) were used in combination with daratumumab (anti-CD38) (online supplemental figure S4A–C). We also studied phagocytosis in an alternative live-cell imaging-based phagocytosis assay that relied on pHrodo-labeled tumor cells turning bright red on phagocytosis (movie in online supplemental files 7 and 8). The advantage of this method was that no counterstain (anti-CD19) was required to determine phagocytic uptake of the target cell. We again tested daratumumab-induced phagocytosis of Daudi cells and found that BYON4228 strongly enhanced uptake of target cells (online supplemental figure S4D–F). The overall potency of BYON4228 (i.e., geometric EC_{50} for all donors, irrespective of the SIRPα genotype) was 0.9 µg/mL in the live-cell imaging pHrodo ADCP assay (N=16) compared with 0.09 µg/mL in the confocal ADCP assay (N=12) (online supplemental figure S4), suggesting the live-cell imaging assay was less sensitive, but was a reproducible orthogonal assay nevertheless. Using the live-cell imaging assay we furthermore studied ADCP of the CRC cell line HT-29 and found that BYON4228 was able to enhance panitumumab-induced and cetuximab-induced phagocytosis (online supplemental figure S5). Altogether, BYON4228 showed functional activity in both neutrophil ADCC and macrophage ADCP assays, indicating its ability to enhance tumor cell killing by myeloid cells.

The BYON4228 silenced Fc tail avoids opsonization and destruction of SIRPα-expressing myeloid cells

With respect to the therapeutic activity of anti-SIRPα antibodies, which relies on anticancer activity of SIRPα-expressing myeloid immune effector cells, it would be undesirable if such SIRPα-expressing cells were depleted through anti-SIRPα Fc mediated processes like CDC, ADCC or ADCP in vivo. Such depletion might compromise both the safety and efficacy of the therapeutic anti-SIRPα antibody. To prevent this, BYON4228 was designed with an IgG1 Fc tail containing L234A/L235A mutations in the Fc tail (abbreviated as ‘LALA’-tail). As expected, BYON4228 displayed strongly decreased binding to FcγRs when compared with its counterpart with wild type (WT) IgG1 backbone without the LALA mutations (IgG1) (online supplemental table S2). Consistent with this, BYON4228 was unable to induce macrophage-mediated phagocytosis of the SIRPα-expressing acute myeloid leukemia (AML) tumor cell line OCI-AML2, whereas the WT IgG1 variant (BYON5664) and anti-CD47 WT IgG1 clearly induced phagocytosis (figure 6A–C). Notably, SIRPAB-11, which contains an IgG1-tail with the K322A modification (designed for reduction of complement activation), also induced ADCP of the OCI-AML2 cells. In line with its inability to induce phagocytosis of SIRPα-positive cells, BYON4228 did not induce NK cell-mediated ADCC of SIRPα-expressing target cells, whereas NK cell induced ADCC was observed by the IgG1 variant (online supplemental figure S6A). Both BYON4228 and its IgG1 variant did not induce CDC of the SIRPα-expressing cell lines U937 and MOLM-13, whereas anti-CD47 (IgG1) induced potent lysis of these cells after incubation with active serum (online supplemental figure S6B). Overall, these data show that the silent Fc tail of BYON4228 prevents destruction of SIRPα-expressing cells by different immune-mediated mechanisms.

Optimal enhancement of anti-TAA-induced ADCC and ADCP by BYON4228 is independent of its Fc functionality

For optimal efficacy, it was suggested previously that the Fc tail of SIRPα-blocking antibodies should not be able to bind to FcγRs on myeloid effector cells in cis to avoid competition with the tumor-opsonizing antibody-FcγR interaction and hence reduced tumor cell destruction, a phenomenon also known as the ‘scorpion effect’.51 59 To examine the influence of the Fc tail for BYON4228
Figure 6  Potent and efficacious enhancement of anti-TAA-induced ADCC and ADCP with BYON4228 having a hIgG1-L234A/L235A Fc tail. (A–C) Macrophage mediated ADCP measured using the live-cell imaging pHrodo ADCP assay after incubation of target cells (OCI-AML2) and effector cells (macrophages) in the presence of a concentration range of indicated antibodies or respective isotype controls. BYON5664 contains the variable domains of BYON4228 but has a wildtype IgG1 constant domain with wildtype effector functions. Donors tested: Two SIRPαBIT/αBIT, three SIRPαBIT/α1, and one SIRPα1/α1 donors. (A) Results show % phagocytosed tumor cells by macrophages from a representative donor. No mAb=effector+target cells only. (B) ADCP EC50 values of all donors. No EC50 values could be calculated from donors with incomplete curve saturation, no response or incorrect curve-fitting as indicated in the graph. (C) The ADCP fold enhancement of all donors. The means are depicted. The fold enhancement=(% phagocytosed cells at 31.6 µg/mL anti-SIRP)/(% phagocytosed cells at effector+target). (D–F) Neutrophil-mediated ADCC measured using the Cr-51 release assay after 4 hours incubation of target cells (SK-BR-3) and effector cells (primary granulocyte-macrophage colony-stimulating factor activated neutrophils) in the presence of a fixed dose of trastuzumab (10 µg/mL) and a concentration range of indicated antibodies or respective isotype controls. BYON5306 is the F(ab’)2 fragment of BYON4228. Donors tested: four SIRPαBIT/αBIT, four SIRPαBIT/α1, and four SIRPα1/α1 donors. (D) Results show % killing of SK-BR-3 cells by activated neutrophils of representative donors with indicated SIRPα genotypes. (E) SIRP-mAb or -F(ab’)2 induced ADCC EC50 values of all donors. No EC50 values could be calculated from donors with incomplete curve saturation, no response or incorrect curve-fitting as indicated in the graph. (F) SIRP-mAb induced ADCC fold enhancement of all donors. The means are depicted. The fold enhancement=(% killing at 10 µg/mL trastuzumab+67 nM anti-SIRPα mAb)/(% killing at 10 µg/mL trastuzumab). For C and F, *p<0.05, **p<0.01, ***p<0.001; p>0.05 is not indicated. P values were calculated by one-way analysis of variance (ANOVA) followed by Tukey’s multiple comparisons test. ADCC, antibody-dependent cellular cytotoxicity; ADCP, antibody-dependent cellular phagocytosis; AML, acute myeloid leukemia; mAb, monoclonal antibody; SIRP, signal-regulatory protein; TAA, tumor-associated antigen.
for optimal in vitro ADCC and ADCP, we generated a fragment antigen-binding region (F(\(\text{ab}^\prime\))\(_{2}\)) version of BYON4228, which was still able to block CD47-SIRP\(\alpha\) axis signaling to the same extent as intact BYON4228 (online supplemental figure S6C). We performed similar side-by-side comparisons to determine enhancement of anti-CAA-induced neutrophil ADCC. BYON4228 was able to enhance trastuzumab-induced ADCC of SK-BR-3 cells with a similar potency and efficacy when the Fc tail was lacking (figure 6D–F). Even when the Fc tail of BYON4228 was replaced with a human WT IgG1 tail (IgG1), ADCC potency and efficacy of enhancement were similar (figure 6D–F). Similar results were also obtained when the BYON4228 variants were tested for enhancement of cetuximab-induced ADCC of A431 cells (online supplemental figure S6D–F). The F(\(\text{ab}^\prime\))\(_{2}\) fragment of BYON4228 was compared with BYON4228 in the rituximab-induced macrophage ADCP assay. Here again, the ability to enhance rituximab-induced phagocytosis of Raji cells was essentially indistinguishable from that of BYON4228 (online supplemental figure S7A–C). Finally, daratumumab-induced ADCP enhancement of Daudi cells was identical for BYON4228 or its F(\(\text{ab}^\prime\))\(_{2}\) (online supplemental figure S7D–F). Overall, these data show that neither BYON4228 nor its WT IgG1 variant suffer from side comparisons to determine enhancement of antibody-dependent tumor cell destruction by both neutrophils and macrophages.

**BYON4228 improves the efficacy of rituximab in human SIRP\(\alpha\)\_scid mice in vivo**

Like most anti-human SIRP\(\alpha\) blocking antibodies, BYON4228 does not cross-react with mouse SIRP\(\alpha\) (data not shown). To study efficacy of BYON4228 in vivo, we generated mice that expressed human SIRP\(\alpha\)\_scid from the Rosa26 promotor only in myeloid cells due to Cre-mediated excision of a STOP cassette (Rosa26-stop\(^{\text{fox}}\)human SIRP\(\alpha\)\_scid) \(\times\) Cebpa\(^{\text{Cre}+/+}\), named huSIRP\(\alpha\)\_scid mice. Western blot analysis confirmed the presence of the human SIRP\(\alpha\) glycoprotein in Cebpa\(^{\text{Cre}+/+}\) mice in conjunction with the endogenous mouse SIRP\(\alpha\) (online supplemental figure S8A). Flow cytometry analysis showed that the mice indeed expressed human SIRP\(\alpha\)\_scid on monocytes and neutrophils (online supplemental figure S8B,C). Furthermore, murine neutrophils that expressed human SIRP\(\alpha\)\_scid enhanced trastuzumab-induced ADCC of SK-BR-3 cells in presence of an anti-SIRP\(\alpha\) antibody (online supplemental figure S8D,E), demonstrating proper functionality of the huSIRP\(\alpha\)\_scid transgene in mice.

First, BYON4228 pharmacokinetic (PK) studies were performed in C57BL/6 mice and in huSIRP\(\alpha\)\_scid mice. In C57BL/6 mice, a low clearance of BYON4228 was noted, while in huSIRP\(\alpha\)\_scid mice, a threefold higher clearance (based on area under the curve (AUC) (\(\text{last}\)) was observed after intravenous or IP administration of a low dose (3 mg/kg) BYON4228 (online supplemental figure S8F,G and online supplemental table S3 and S4). This difference in drug clearance is most likely due to target-mediated drug disposition (TMDD). In huSIRP\(\alpha\)\_scid mice, the drug elimination via TMDD could be saturated by high doses (intravenous) or repeat dosing (IP) of BYON4228, leading to prolonged high plasma BYON4228 levels (online supplemental figure S8F–H).

Next, antitumor efficacy of rituximab alone or in combination with BYON4228 was studied in huSIRP\(\alpha\)\_scid mice xenografted with Raji tumors. No significant effects of BYON4228 alone on tumor growth were observed (one experiment, N=9 mice/group, data not shown). Rituximab treatment at 1 mg/kg significantly inhibited growth of Raji cells in huSIRP\(\alpha\)\_scid mice and this effect could be enhanced by combination treatment with BYON4228 at 5 mg/kg (figure 7A), one of two independent studies shown, N=11 mice/group). These results show that the huSIRP\(\alpha\)\_scid mouse model can be employed to study efficacy of anti-SIRP\(\alpha\) antibodies in mouse. Western blot analysis confirmed the presence of the human SIRP\(\alpha\) glycoprotein in Cebpa\(^{\text{Cre}+/+}\) mice in conjunction with the endogenous mouse SIRP\(\alpha\) (online supplemental figure S8A). Flow cytometry analysis showed that the mice indeed expressed human SIRP\(\alpha\)\_scid on monocytes and neutrophils (online supplemental figure S8B,C). Furthermore, murine neutrophils that expressed human SIRP\(\alpha\)\_scid enhanced trastuzumab-induced ADCC of SK-BR-3 cells in presence of an anti-SIRP\(\alpha\) antibody (online supplemental figure S8D,E), demonstrating proper functionality of the huSIRP\(\alpha\)\_scid transgene in mice.

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**Figure 7** BYON4228 potentiates the antitumor activity of rituximab in vivo and pharmacokinetic evaluation of BYON4228 in cynomolgus monkeys. (A) Growth curve of Raji cells (subcutaneous) in female huSIRP\(\alpha\)\_scid mice treated IP with vehicle, rituximab monotherapy, or rituximab in combination with BYON4228. All compounds were administered IP three times a week (indicated by arrows between dotted lines). Rituximab was used at 1 mg/kg for monotherapy and in combination with 5 mg/kg BYON4228. N=11 per group. Statistical analysis of area under the curve from randomization to end of study: *** \(p<0.001\), Dunnett’s multiple comparison test. (B) Mean BYON4228 plasma concentrations in cynomolgus monkey after a single intravenous infusion of BYON4228 (N=2 (one female and one male) ±SD/dose group). IP, intraperitoneal; SIRP, signal-regulatory protein; scid, severe combined immunodeficiency.
vivo and, most importantly, that BYON4228 can improve tumor clearance when combined with rituximab. These data are in line with Murata et al, who demonstrated enhancement of the inhibitory effect of rituximab by anti-human SIRPα antibody on the growth of tumors formed by Raji cells in other human SIRPα transgenic mice. 61

BYON4228 has a favorable preclinical safety profile

We assessed the safety of BYON4228 in vitro. BYON4228 did not display hemolytic activity, it did not induce red blood cell (RBC) clumping and it was compatible with human plasma in vitro (data not shown). At concentrations of 1 up to 100 µg/mL, BYON4228 did not induce any biologically significant increases in a panel of 18 cytokines/chemokines measured (data not shown). A toxicological evaluation of BYON4228 was performed in cynomolgus monkeys. We noticed that BYON4228 displayed potent binding to SIRPα-positive granulocytes of most cynomolgus monkeys (online supplemental figure S9A,B). Individual monkeys that displayed no or very poor BYON4228 binding were deselected for toxicity studies. The selected individuals had an average 3.5-fold more potent binding (3.5-fold lower EC50) compared with that in man (0.03 µg/mL, online supplemental figure S9B vs 0.09 µg/mL, figure 1D), supporting the relevance of cynomolgus monkey for the human risk assessment of BYON4228. Intravenous infusion of BYON4228 was well tolerated as a single dose up to 100 mg/kg. PK profiles in monkeys showed almost dose linear increase in exposure. One week after dosing, variability is observed in the PK curves due to the induction of immunogenicity as was confirmed by the observation of anti-drug antibodies (figure 7B, online supplemental table S5 and data not shown). In the 5-cycle repeated dose toxicity study in monkey, where 3, 10 or 30 mg/kg BYON4228 was infused intravenously one time a week, BYON4228 showed comparable PK profiles and was well tolerated with no adverse effects on body weight and no clinical evidence of toxicity including safety pharmacology endpoints (cardiovascular, respiratory, and central nervous system) (data not shown). In addition, no effects on hemoglobin levels, RBC count or platelets were noted (online supplemental figure S9C–E). When comparing the measured SIRPα receptor occupancy (RO) data with the observed exposure levels, the data show that >80%–90% RO (needed for 100% efficacy in ADCC and ADCP assays in vitro, data not shown) can be reached at plasma concentrations around 2 µg/mL. BYON4228 based on limited data. The mean CLOUGH of BYON4228 following the first dose in the 5-cycle toxicity study ranged from 15.8 µg/mL to 285 µg/mL for the 3–30 mg/kg doses indicating that the >80% RO had been reached over the entire dosing interval. Based on the 5-cycle study, the No Adverse Effect Level (NOAEL) was considered to be the highest dose tested, 30 mg/kg. At this repeated dose, the exposure is 165,000 hours µg/mL for males and 179,000 hours µg/mL for females.

DISCUSSION/CONCLUSION

The CD47-SIRPα myeloid immune checkpoint has been identified as a promising therapeutic target to promote tumor destruction in combination with anti-TAAAs and other therapeutics. 9 38 Blockade of CD47-SIRPα signaling can stimulate destruction of antibody-opsonized cancer cells by innate immune cells, in particular by macrophage ADCC and neutrophil ADCC. Furthermore, disruption of CD47-SIRPα interactions triggers adaptive T cell-mediated anticancer immunity, probably, at least in part, by enhancing cross-presentation by SIRPα-expressing dendritic cells. Currently, a number of agents targeting CD47 or SIRPα are in clinical development. The vast majority of these agents are targeting CD47 and the most advanced ones have already shown encouraging safety/efficacy profiles in early clinical studies. 36 37 39 41 Nevertheless, as outlined above there are a number of disadvantages of targeting CD47, and this may therefore not be the most optimal and selective way to antagonize CD47-SIRPα interactions therapeutically. Notably, the disruption of CD47-SIRPγ interactions, which are critical for T-cell extravasation and activation, 34 46–48 might curtail durable adaptive antitumor immunity.

As an alternative strategy, therapeutic agents have been developed that target SIRPα 29 34 49–55 and at present four of such drugs are, or have been, in early clinical development. Preclinical data of HEFLB, SIRPAB-11, and 1H9, most likely corresponding to BI 765063 (formerly OSE-172), BMS-986351 (anzurstobart, formerly CC-95251), and GS-0189 (formerly FSI-189) (table 1), respectively, have been reported. 34 49–58 However, we have shown here that these SIRPα blocking antibodies, like several others that have been reported, 50–53 35 either fail to recognize both of the two SIRPα polymorphic variants present in the human population, or they cross-react with SIRPγ.

Here we preclinically characterize BYON4228, a novel anti-SIRPα antibody, and perform direct in vitro comparisons with HEFLB, SIRPAB-11, and 1H9. Our results identify BYON4228 as a highly differentiating antibody, which recognizes both of the two polymorphic variants SIRPαHET and SIRPα1. BYON4228 efficiently inhibits CD47 binding to primary myeloid cells from individuals with all three genotypes (ie, SIRPαHET-homozygotes and SIRPα1-homozygotes and SIRPαHET/SIRPα1-heterozygotes). The antagonistic activity of BYON4228 can be explained by direct competition with CD47 binding, as the BYON4228 epitope strongly overlaps with the previously described CD47 binding site on both of the SIRPα variants. 21 In line with this, BYON4228 prevents inhibitory signaling via the CD47-SIRPα axis, and promotes neutrophil ADCC and macrophage ADCP using immune effector cells from individuals of all three SIRPα genotypes. In contrast, HEFLB lacks binding to SIRPα1, and, as a consequence, the ability to reduce CD47 binding to SIRPα, homozygous myeloid cells, and clearly fails to enhance ADCC and ADCP by SIRPα, homozygous myeloid immune effector cells. Notably, HEFLB also appears to be only a partial inhibitor of CD47 binding by SIRPαHET/SIRPα1.
heterozygote myeloid cells, and, if anything, only a weak stimulator of ADCC and ADCP, mediated by, respectively, neutrophils and macrophages from such heterozygous individuals. These findings are also supported by previous reports.\textsuperscript{31} \textsuperscript{35} These properties will likewise compromise the applicability of BI 765063/OSE-172, the probable corresponding agent of HEFLB in clinical development, described not to bind to T cells,\textsuperscript{49} only a narrow window to inhibit CD47 binding to primary T cells. While 1H9 was active putative clinical equivalents BMS-189 (NCT04502706) was terminated after inclusion of a patient with anemia and thrombocytopenia. The latter contrasts with results obtained for several of the more advanced CD47-blocking antibodies, including magrolimab, in both preclinical toxicity studies and clinical studies, where anemia and thrombocytopenia were clearly observed.\textsuperscript{36} \textsuperscript{39} \textsuperscript{66} \textsuperscript{67} Collectively, these data show that BYON4228 is a highly differentiating pan-allelic blocking anti-SIRP\textgreek{a} antibody, which specifically lacks reactivity with SIRP\textgreek{a} on T cells. In doing so, BYON4228 has the principal features to potentially become a best-in-class drug. Our preclinical functional ADCC and/or ADCP experiments with a variety of target cancer cells and clinically available therapeutic anti-TAs, including trastuzumab, cetuximab, panitumumab, rituximab and daratumumab, underline the broad potential applicability of BYON4228. This may also extend to other non-anti-TA combinations, such as other checkpoint inhibitors\textsuperscript{33} \textsuperscript{34} \textsuperscript{41} \textsuperscript{49} chemotherapy\textsuperscript{68} and radiotherapy.\textsuperscript{65} First-in-human clinical studies with BYON4228 are scheduled to start in 2023.
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Contributors MJvH and TkvdB wrote the manuscript with input from all authors. MJvH, SAZ, RJA, HLM, KF, EM-H, DRFfG, DEJWvW, JS, WAK, DvdD, MT, RU, GJAR, GV, MMcvDL, WHAD and TKvdB contributed to the conception and development of the project. SAZ, IMJR-B, MBcpB, LD-E, KdL-A, DD, EWHS-L, DWJk, IL, HO, HLM, KF, EM-H, MEMS, PBd and Bwd performed experiments and analyzed data. DRFfG, DEJWW, Lj-J and JS generated antibodies (fragments), recombinant proteins and cell lines. RJA, NvdWel, JWHS, WAK, DvdD and RU coordinated in vivo studies and (PK) analysis. All authors read and approved the final version of the manuscript. MJvH is responsible for the overall content as guarantor.

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