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Identification and characterisation of carnostatine (SAN9812), a potent and selective carnosinase (CN1) inhibitor with in vivo activity

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Abstract
Carnosinase 1 (CN1) has been postulated to be a susceptibility factor for developing diabetic nephropathy (DN). Although its major substrate, carnosine, is beneficial in rodent models of DN, translation of these findings to humans has been hampered by high CN1 activity in human serum resulting in rapid degradation of carnosine. To overcome this hurdle, we screened a protease-directed small-molecule library for inhibitors of human recombinant CN1. We identified SAN9812 as a potent and highly selective inhibitor of CN1 activity with a $K_i$ of 11 nM. It also inhibited CN1 activity in human serum and serum of transgenic mice-overexpressing human CN1. Subcutaneous administration of 30 mg/kg SAN9812 led to a sustained reduction in circulating CN1 activity in human CN1 transgenic (TG) mice. Simultaneous administration of carnosine and SAN9812 increased carnosine levels in plasma and kidney by up to 100-fold compared to treatment-naïve CN1-overexpressing mice. To our knowledge, this is the first study reporting on a potent and selective CN1 inhibitor with in vivo activity. SAN9812, also called carnostatine, may be used to increase renal carnosine concentration as a potential therapeutic modality for renal diseases linked to glycoxidative conditions.

Keywords Diabetic nephropathies · Carnosine · Aminoacyl-histidine dipeptidase · Drug discovery

Introduction
The rising global prevalence of type 2 diabetes (T2D) is accompanied by an increase in diabetic complications, such as diabetic nephropathy (DN), the single most common cause of end-stage renal disease (ESRD). Despite our continuously evolving understanding of the pathogenesis of DN and potential interventions, the current standard of care has largely failed to achieve stable long-term results, and a...
strong unmet need for novel and more efficacious therapies remains (de Zeeuw and Heerspink 2016).

Carnosine (β-alanyl-l-histidine) is a physiological dipeptide first discovered in meat extract more than 100 years ago (Gulewitsch and Amiradžibi 1900). It is primarily found in skeletal muscle where its content can be as high as 2 g/kg wet tissue or 0.2% of overall skeletal muscle mass (Boldyrev et al. 2013). It is also present in other tissues such as brain, kidney or spleen albeit at 10- to 100-fold lower concentrations (Kamal et al. 2009). Properties pertaining to the physiological roles of carnosine include its pH-buffering capacity, scavenging of reactive oxygen species (ROS) and peroxyl radicals, metal-ion chelation and protection against advanced glycation and lipoxidation end products (AGE and ALE, Boldyrev et al. 2013). Notably, carnosine has also been demonstrated to reverse pre-formed glycated proteins (Seidler et al. 2004). In line with its role in muscle physiology, food supplements containing carnosine or its precursor beta-alanine improve exercise performance in untrained individuals (Hill et al. 2007) as well as in elite athletes (Van Thienen et al. 2009; Baguet et al. 2010). Due to its metal-ion chelating and anti-oxidative properties, carnosine can ameliorate transition metal-catalysed- or hyperglycaemia-mediated oxidative stress which may occur in several (pathologic) conditions such as diabetes and its complications (Zhang et al. 2016). Apart from the nutritional intake, tissue carnosine concentration also depends on the activity of local carnosine synthase, an ATP-dependent enzyme, as shown for a variety of tissues, e.g., skeletal muscle, brain or kidney (Drozak et al. 2010; Peters et al. 2015).

Carnosinase (CN1, EC 3.4.13.20) is the major carnosine-degrading enzyme. The human gene was cloned in 2003 and identified as the gene encoding the secreted form of carnosinase (Teufel et al. 2003). By positional cloning in a large Turkish family affected by DN, the carnosine dipeptidase 1 gene (CNDP1) was identified as a susceptibility gene for developing DN (Vardarli et al. 2002; Janssen et al. 2005). The leucine repeat in the signal sequence of CNDP1 gene is essential for CN1 secretion with the short allelic variant (CTG)5 associated with low-serum CN1 concentrations. Importantly, patients with T2D without DN are significantly more frequently homozygous for the (CTG)5 allele (Freedman et al. 2007; Riedl et al. 2007; Alkhalf et al. 2010; Peters et al. 2016). It was postulated that low CN1 activity might favour high circulating carnosine concentrations. Of note, in a carnosine or beta-alanine supplementation study in rats with diet-induced obesity, it was recently demonstrated that plasma rather than muscle carnosine is involved in ameliorating high-fat diet-induced lipoxidative and inflammatory stress (Stegen et al. 2015). Carnosine supplementation studies in murine models demonstrated therapeutic efficacy of carnosine to ameliorate diabetes and DN, as albuminuria and histological lesions were reduced after supplementation (Lee et al. 2005; Alhamdani et al. 2007; Sauerböer et al. 2007; Riedl et al. 2011; Aldini et al. 2011; Nagai et al. 2012; Peters et al. 2012; Albrecht et al. 2017).

However, rodents—with the exception of Syrian hamster—lack the signal peptide in the CN1 gene and consequently do not have CN1 in the circulation (Boldyrev et al. 2013). This results in increased carnosine levels after oral carnosine supplementation (Albrecht et al. 2017). In contrast, carnosine supplementation in humans leads only to a small increase in circulating carnosine because of its fast degradation by serum CN1 (Everaert et al. 2011). Because of this difference, extrapolation of rodent findings to humans is difficult. As T2D patients have even higher serum CN1 activity levels compared to healthy controls (Riedl et al. 2010), CN1 inhibition could be a potential strategy to increase the serum concentration of carnosine, especially in combination with carnosine supplementation. So far, this possibility has not been explored due to the lack of a specific CN1 inhibitor. Here, we describe the identification and characterisation of a potent and selective CN1 inhibitor, carnostatine, and its effect on serum CN1 activity and carnosine levels in serum and renal tissue after co-administration with carnosine.

Results

We screened a protease-directed small-molecule library containing 6080 compounds for inhibitors of human recombinant CN1 using a screening assay based on the detection of histidine generated by proteolytic cleavage of carnosine (supplementary methods). The human recombinant CNDP1 showed a \( K_m \) value of 190 \( \mu \)M (supplementary figure 1), similar to the previously reported value of 175 \( \mu \)M (Peters et al. 2010). The average \( Z' \) value (± SD) of all 38 test plates was \( 0.85 ± 0.06 \) (supplementary figure 2), indicating that the assay was of excellent robustness (Zhang et al. 1999). Results are summarised in Fig. 1. In total, we identified 133 compounds with > 50% inhibition. These compounds were re-tested in duplicates (supplementary figure 3).

One of the most potent compounds identified was SAN9812 (Fig. 2), with an \( IC_{50} \) value of 18 \( nM \) on human recombinant CN1 at a carnosine concentration of 200 \( \mu \)M, i.e., close to the \( K_m \) of 190 \( \mu \)M. SAN9812 was found to be a competitive inhibitor, showing a right-shift in its \( IC_{50} \) value with increasing carnosine concentration (Fig. 3a). The \( K_i \) value of SAN9812 was determined as 11 \( nM \) (Fig. 3b). The synthesis of SAN9812 is described in supplementary figure 4 and supplementary methods.

The inhibitory activity of SAN9812 on human CN1 was confirmed in serum from TG mice-expressing human CN1 and serum from healthy human volunteers (Fig. 3c, d). \( IC_{50} \) values were determined as 650 and 340 \( nM \), respectively, at a carnosine concentration of 0.2 mg/ml or
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880 µM. The average baseline hCN1 activities in transgenic mice and healthy human volunteers were 137 ± 25 and 1.1 ± 0.25 µmol/ml/h, respectively.

Of note, SAN9812 was found to be inactive on a standard panel of 33 different receptors, channels, transporters and enzymes (supplementary Table 1) when tested at a concentration of 10 µM, reflecting overall selectivity of SAN9812 and ruling out that it is a modulator of a broad spectrum of biological activities. The ADME/PK (absorption, distribution, metabolism, excretion and pharmacokinetics) profile of SAN9812 is summarised in Table 1. The compound is very stable in human, rat and mouse liver microsomes and shows moderate clearance in primary human hepatocytes. Permeability in the CaCo-2 assay as a measurement of intestinal permeability was determined as 17.9 nm/s, indicating moderate oral absorption potential.

Plasma levels of SAN9812 after single bolus administration of either 3 mg/kg i.v. or 30 mg/kg s.c. are shown in Fig. 4a. Clearance, volume of distribution and subcutaneous bioavailability are summarised in Table 1. After subcutaneous injection, maximal plasma concentrations were achieved already at the first sampling time point after administration (15 min). The maximal compound concentration after s.c. administration was 80 µg/ml, corresponding to approximately 300 µM. Plasma levels quickly decrease to 30 µg/ml (120 µM) after 30 min, 5.7 µg/ml (22 µM) after 1 h, and 350 ng/ml (1.4 µM) after 2 h. The lower level of quantification for the detection method was 10 ng/ml (40 nM). After intravenous injection, maximal plasma concentration was also observed at the first sampling time point, but there was a fast decrease in plasma SAN9812 concentrations reflecting rapid clearance of the compound from the circulation. Further experiments were performed using subcutaneous injection due to the longer availability of the compound in the blood.

In a subsequent pharmacodynamics study, CN1 activity was measured after a single subcutaneous bolus of 30 mg/kg SAN9812, at the same time points as previously in the PK experiment. Results are shown in Fig. 4b, c. Animals were stratified according to their baseline serum carnosinase activity into a high-activity- (> 250 µmol/ml/h) and a low-activity (< 45 µmol/ml/h) group. Mean baseline activities in the two groups were 259 ± 13 and 7.4 ± 4.9 µmol/ml/h, respectively. Administration of SAN9812 led to a rapid reduction in serum carnosinase activity by more than 90% in both groups which was sustained for > 8 h. At the last sampling time point 16 h after compound administration, serum carnosinase activity was still reduced by 40% in the low-activity group and by 75% in the high-activity group.

SAN9812 was then subcutaneously administered over 7 days once daily at a dose of 30 mg/kg. Blood samples were taken on days 1, 4 and 7 before, 6 and 12 h after the injection to measure CN1 activity. The results are depicted in Fig. 5. Again, CN1 activity was strongly and rapidly reduced after the first SAN9812 injection on day 1 and remained low over a period of > 12 h. Importantly, baseline CN1 activities on days 4 and 7, measured 24 h after the previous SAN9812 administration on days 3 and 6, respectively, were considerably lower compared to CN1 activity on day 1 before the
first injection, indicating sustained carnosinase inhibition over 24 h in both the low-activity and high-activity groups.

The same average inhibition in percent was achieved in the group with low-baseline activity as in the group with high-baseline activity. In mice with low-baseline activity, CN1 activity was efficaciously inhibited over the whole observation period with an activity level kept below 10 µmol/ml/h. In the group with high-baseline activity levels, absolute CN1 activity could sometimes exceed 100 µmol/ml/h between injections, although percent-inhibition values were comparable in all animals (Fig. 5).

We did not note any obvious adverse effects of treatment with SAN9812 over 7 days, body weight and blood glucose were not different between treated and untreated animals.

To determine whether serum CN1 inhibition by SAN9812 can increase carnosine concentration in vivo in hCN1 overexpressing mice, animals were given a single bolus of 200 mg carnosine alone or in combination with 30 mg/kg SAN9812 per s.c. injection. Plasma samples were taken 4 and 8 h after injection. Kidneys were harvested 8 h after injection. CN1-negative wild type mice and hCN1 mice without treatment served as controls.

Compared to wild type animals, hCN1 TG mice had significantly lower plasma carnosine levels (Fig. 6a, b). A similar, but non-significant trend was also seen for the kidney (Fig. 6c). As shown in Fig. 6a, in hCN1 mice after a single

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**Table 1** ADME and PK profile of SAN9812

| Metabolic stability h/r/m microsomes (% remaining) | 100/94/99 |
| Human hepatocyte clearance (ml/h/10⁶ cells) | 0.032 |
| Caco-2 permeability (nm/s) | 17.9 |
| Clearance (mL/min/kg) | 0.344 |
| Volume of distribution (L/kg) | 0.122 |
| Subcutaneous bioavailability (%) | 42.5 |
| Selectivity panel | Inactive

*At a concentration of 10 µM vs a panel of 33 diverse receptors, ion channels, transporters and enzymes*

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**Fig. 3** a Concentration–response curves for SAN9812 determined at 100 µM (open triangles) and 1000 µM (filled squares) carnosine with human recombinant CN1 in PBS (n=6 each), corresponding IC₅₀ values were determined as 10 and 54 nM, respectively. b Dependence of SAN9812 IC₅₀ values on carnosine concentration (n=6 each). The Kᵢ values defined as the intersection of the linear regression line (R²=0.91) with the y-axis was determined as 11 nM. c Concentration–response curve of SAN9812 in serum from transgenic mice-expressing human CN1 (three replicates from three mice), and d in human serum (three replicates from two subjects) at a carnosine concentration of 880 µM (0.2 mg/ml). IC₅₀ values were determined as 650 and 340 nM, respectively. Data are presented as mean ± SEM.
subcutaneous bolus of carnosine, the plasma carnosine concentration increased compared to hCN1-overexpressing controls without carnosine administration \((p = 0.049)\) 4 h after injection but was still lower compared to wild type mice without injection \((p = 0.058)\). In contrast, a concurrent administration of carnosine and SAN9812 led to a further increase in carnosine plasma concentration, up to the levels of hCN1-negative controls without injection 4 h after administration. 8 h after concurrent administration of carnosine and SAN9812, plasma carnosine levels had returned to levels of hCN1 controls probably due to residual CN1 activity \((\text{mean} \pm \text{SD} 14.4 \pm 15.7 \, \mu\text{mol/ml/h})\). Mean CN1 activity in hCN1 mice was 55.3 ± 30.6 \mu\text{mol/ml/h}.

As shown in Fig. 6c, in the kidney concurrent administration of carnosine and SAN9812 significantly increased carnosine concentration \((\text{mean} \pm \text{SD} 3.5 \pm 2.3 \, \text{nmol/mg})\) compared to wild type controls without any treatment \((0.32 \pm 0.46 \, \text{nmol/mg})\), hCN1-overexpressing mice \((0.034 \pm 0.028 \, \text{nmol/mg})\) and also hCN1-overexpressing mice supplemented with carnosine \((0.52 \pm 1.2) \, (p < 0.005)\). Of note, kidneys were removed 8 h after injection when plasma carnosine levels had already returned to normal.

Discussion

In this work, we discovered carnostatine (SAN9812) as a potent and selective CN1 inhibitor with a \(K_i\) of 11 nM for human recombinant CN1. Subcutaneous injection of SAN9812 led to a sustained reduction in serum carnosinase activity in transgenic mice overexpressing human CN1, both acutely as well as upon subchronic treatment. Additionally, a combination of carnosine and carnostatine caused a further increase in serum carnosine concentration and, for a disease such as DN most importantly, an increase of carnosine levels in renal tissue.

Carnosinase 1 has attracted considerable interest through the discovery in humans that a polymorphism of CN1 associated with lower serum CN1 concentrations protects against DN. In keeping with the hypothesis that low serum CN1 concentration may protect patients with type 2 diabetes from developing DN, the rationale of the current study was to generate a potent CN1 inhibitor as a potential drug candidate for preventing or slowing down the progression of DN.

The protective effect of a genotype with lower CN1 activity, as shown in large cross-sectional studies, is likely...
Fig. 5 Subchronic (7 days) administration of SAN9812 (30 mg/kg s.c. q.d.) inhibits serum CN1 activity. a Serum CN1 activity (relative to day 1 baseline) prior to, 6 and 12 h after compound administration on days 1, 4 or 7 for all mice investigated (n = 10, mean ± SEM). b, c Individual absolute CN1 activities for the high-activity (b, n = 5) and the low-activity (c, n = 5) group of hCN1-expressing transgenic mice.

Fig. 6 Plasma and renal tissue carnosine concentration upon subcutaneous carnosine supplementation (CAR) with and without SAN9812 administration in hCN1 overexpressing (CN1+) mice (n = 5–8 per group). hCN1-negative wild type mice (CN1−) served as controls. The lower level of quantification for the detection method was 0.0001 nmol/mg. Data are shown as mean ± SD. a Plasma carnosine concentration 4 h and b 8 h after administration of carnosine ± SAN9812. c Carnosine concentration in the kidney harvested 8 h after injection. One-way ANOVA followed by Dunnett’s multiple comparisons test was performed using GraphPad Prism (*p < 0.05, ** for p < 0.01)
mediated by an increase in serum carnosine concentration. To increase serum carnosine, many carnosine supplementation studies have been conducted in rodent models. However, as rodents lack the signal peptide in the CN1 gene and consequently do not have CN1 in the circulation (Boldyrev et al. 2013), the translation of these findings to humans is difficult. Unlike rodents, humans have considerable serum CN1 activity which may even be higher in particular disease states, e.g., in diabetic individuals compared to healthy controls (Riedl et al. 2010). Thus, as observed in healthy volunteers, carnosine supplementation only leads to a moderate increase in plasma carnosine for a limited period of time between 1 and 2 h because of its rapid degradation by CN1 (Everaert et al. 2012) that may markedly limit carnosine’s beneficial effects in humans.

Several human trials in the field of sports medicine have been conducted, either with carnosine or beta-alanine, the rate-limiting precursor for endogenous carnosine synthesis, and showed beneficial effects on exercise performance (Hill et al. 2007; Van Thienen et al. 2009; Baguet et al. 2010). Although muscle carnosine was not measured in our study, the marked increase by 100-fold in renal tissue concentration of carnosine by simultaneous administration of carnosine and a CN1-specific inhibitor, although in an artificial setting, was promising and suggests that our approach may also enhance the beneficial effects of carnosine in other tissues.

Other human trials have shown a positive effect of carnosine supplementation on insulin sensitivity in obese human individuals (De Courten et al. 2016; Regazzoni et al. 2016). Because of the reasons given above, an additional CN1 inhibition may further increase these beneficial effects. So far, this has not been studied because of the lack of a specific CN1 inhibitor. Here, we describe SAN9812, a specific CN1 inhibitor as a new tool for such studies. As SAN9812 can stabilise carnosine levels both in plasma and kidney tissue after co-administration with carnosine, we propose to use the name carnostatine for SAN9812.

In the current work, carnostatine and carnosine were subcutaneously injected. The oral availability of carnostatine has so far not been investigated. Oral absorption potential of carnostatine, based on Caco-2 permeability, was found to be moderate but high enough to merit additional studies where the compound could be given via oral gavage or drinking water. Especially the latter option would make it feasible to perform chronic treatment studies with minimal compound application stress for the study animals.

Possible side effects of carnostatine could be similar to carnosinemia, where a lack of carnosinase is associated with several neurological symptoms. Although we could not detect any major side effects after 7 days of carnostatine treatment in mice, we could not exclude any neurological symptoms since no behaviour test was performed. Moreover, CN1 is postulated to play a major role in brain gamma-aminobutyric acid (GABA)-ergic system as it can release GABA by degrading homocarnosine. If carnostatine can bypass the blood–brain-barrier, it could interfere with this system and cause GABA-deficiency symptoms such as depression, anxiety or panic. In addition, pathways involving other degradation products of CN1, e.g., histidine or beta-alanine could be influenced.

The model system used in our study—mice-overexpressing human CN1—is reflecting the situation in humans where plasma carnosinase activity can be detected. However, the model also has certain limitations, such as the high variability of CN1 activity and a high absolute plasma CN1 activity compared to lean human volunteers. Thus, care should be taken to stratify animal according to CN1 activity and to select animals with moderately increased CN1 activity for further chronic studies.

To our knowledge, this is the first in vivo active carnosinase inhibitor described. This CN1 inhibitor is potent with a $K_i$ of 11 nm and specific with no reactivity vs. a panel of 33 different receptors, channel transporters and enzymes. So far, our studies were limited to healthy animals and short treatment periods. It will be interesting to see whether chronic carnosinase inhibition, especially if carnostatine is orally administered in combination with carnosine, can protect from diabetic nephropathy as seen in carnosine supplementation studies in mice devoid of plasma CN1 activity. Such studies are currently underway.

**Methods**

**Subjects (mice, human)**

**Mice**

Human CNDP1 transgenic mice on a BTBR wt/ob background (Jackson Laboratory Stock number: 004824) were used for the experiments. These mice express human CN1 in brain and liver from where it is secreted into the blood. The generation and characterisation of these hCN1 over-expressing mice will be described in a separate paper. The animals were housed in a 12-h light/dark cycle at 22 °C and sacrificed under isoflurane anaesthesia by neck dislocation. Animals between 8 and 10 weeks of age with a body weight of 25–35 g were used. For the experiment with carnosine ± carnostatine injection age- and sex-matched animals were used. Body weight and blood glucose were measured between the injections to detect major abnormalities of animal well-being. No difference between two groups was detected.
Human

Serum samples were taken from two healthy donors (one male, one female) that were 25 and 26 years of age, respectively.

High-throughput screening and compound characterisation

We screened a protease-directed small compound library using human recombinant carnosinase 1 based on an assay described by Teufel 2002. Details are described in supplementary methods. IC$_{50}$ and $K_i$ values and the mode of inhibition by SAN9812 were determined as described in supplementary methods. Activity of SAN9812 on a standard panel of 33 different receptors, channels, transporters and enzymes was measured at Eurofins Pharma Discovery Services. Detailed descriptions of individual assays can be found at https://www.eurofinsdiscoveryservices.com.

Activity of SAN9812 in human and mouse serum

To determine the concentration–response curve of SAN9812, serum from two healthy donors and three CN1-transgenic mice were used. SAN9812 was diluted with PBS and added to the serum directly (2–3 s) before measuring the CN1 activity.

ADME profile of SAN9812

A detailed description of the methods used to investigate the ADME profile can be found in the supplementary methods.

Pharmacokinetics

Pharmacokinetic parameters were determined in female C57BL/6 mice (Harlan, Germany). After a single administration of either 3 mg/kg i.v. or 30 mg/kg s.c. of test compound dissolved in phosphate-buffered saline, pH 7.4, plasma samples were taken at eight time points over 24 h. Three animals per time point were used. Concentrations of the test compound were determined with an exploratory LC–MS/MS method using an internal standard. Clearance and volume of distribution were determined based on the i.v. pharmacokinetics, subcutaneous bioavailability by comparing the area under the curve after subcutaneous vs. intravenous administration. Pharmacokinetic parameters were calculated using the program WinNonlin 6.4 (Pharsight Corporation) assuming a non-compartmental model and linear trapezoidal interpolation.

Pharmacodynamics

The mice are injected with 30 mg/kg SAN9812 in a 0.9% NaCl solution subcutaneously. Before and after the injection blood was taken at different time points. The blood samples were incubated for 30 min at room temperature before being centrifuged at 2500g for 10 min. The supernatant as serum was used afterwards to determine the CN1 activity. Two groups of animals with three animals each were used, one with activity levels all below 45 µmol/(ml h) defined as the low-activity group and one with activity levels all above 250 µmol/(ml h) defined as high-activity group. These two groups are depicted separately in the figures.

Pharmacodynamics upon subchronic administration

The mice were treated with 30 mg/kg SAN9812 in a 0.9% NaCl solution for 1 week by daily subcutaneous injection. Before and after the compound administration serum was taken and CN1 activity was measured. Animals were stratified to high- and low-activity groups as described above and five mice per group were used.

Acute carnosine injection ± SAN9812 in mice

Carnosinase 1-overexpressing mice were supplemented with a bolus of 200 mg carnosine alone or in combination with 30 mg/kg SAN9812 in a 0.9% NaCl solution by s.c. administration. hCN1-negative and hCN1-overexpressing mice served as controls ($n = 5–8$). Plasma samples were taken after 4 h and after 8 h. After 8 h, mice were sacrificed and kidneys were harvested.

Chemical analyses

A more extended description of chemical analyses can be found in the supplementary methods.

Sample preparation

Serum and EDTA plasma were taken according to standard protocols. For CN1, activity serum was taken, for carnosine measurement, EDTA plasma and tissue samples supplemented with 2 mmol EDTA were used.

CN1 activity

Carnosinase 1 activity was measured using a modified protocol based on the assay described by Teufel 2002. Details are in the supplementary methods.
Carnosine concentration

Carnosine concentration was measured by high-performance liquid chromatography as previously described (Schönherr 2002).

Figures and statistics

Figures were created using Microsoft Excel and GraphPad Prism 7.0. Error bars denote either standard deviation (SD) or standard error of the mean (SEM) as described in the individual figure legends. To compare different groups, ANOVA one-way with Holm-Sidak’s multiple comparison test were applied using GraphPad Prism 7.0.

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Compliance with ethical standards

Conflict of interest

AK., S.R. and M.T. are or were employees of Sanofi Aventis Deutschland GmbH.

Ethical approval

All procedures performed in studies involving animals were in accordance with the ethical standards of the institution or practice at which the studies were conducted. The experiments were approved by the local Ethical Committee.

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