Effects of Inhibiting Dipeptidyl Peptidase-4 (DPP4) in Cows with Subclinical Ketosis

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Abstract

The inhibition of dipeptidyl peptidase-4 (DPP4) via specific inhibitors is known to result in improved glucose tolerance and insulin sensitivity and decreased accumulation of hepatic fat in type II diabetic human patients. The metabolic situation of dairy cows can easily be compared to the status of human diabetes and non-alcoholic fatty liver. For both, insulin sensitivity is reduced, while hepatic fat accumulation increases, characterized by high levels of non-esterified fatty acids (NEFA) and ketone bodies. Therefore, in the present study, a DPP4 inhibitor was employed (BI 14332) for the first time in cows. In a first investigation BI 14332 treatment (intravenous injection at dosages of up to 3 mg/kg body weight) was well tolerated in healthy lactating pluriparous cows (n = 6) with a significant inhibition of DPP4 in plasma and liver. Further testing included primi- and pluriparous lactating cows suffering from subclinical ketosis (β-hydroxybutyrate concentrations in serum > 1.2 mM; n = 12). The intention was to offer effects of DPP4 inhibition during comprehensive lipomobilisation and hepatosteatosis. The cows of subclinical ketosis were evenly allocated to either the treatment group (daily injections, 0.3 mg BI 14332/kg body weight, 7 days) or the control group. Under condition of subclinical ketosis, the impact of DPP4 inhibition via BI 14332 was less, as in particular β-hydroxybutyrate and the hepatic lipid content remained unaffected, but NEFA and triglyceride concentrations were decreased after treatment. Owing to lower NEFA, the revised quantitative insulin sensitivity check index (surrogate marker for insulin sensitivity) increased. Therefore, a positive influence on energy metabolism might be quite probable. Minor impacts on immune-modulating variables were limited to the lymphocyte CD4+/CD8+ ratio for which a trend to decreased values in treated versus control animals was noted. In sum, the DPP4 inhibition in cows did not affect glycaemic control like it is shown in humans, but was able to impact hyperlipemia, as NEFA and TG decreased.
Introduction

Dipeptidyl peptidase-4 (DPP4) plays a major role in glucose metabolism and is responsible for the degradation of incretin hormones, such as glucagon-like peptide-1 (GLP-1). Human GLP-1 is released from the small intestine in response to oral glucose [1]. It stimulates insulin secretion via activating specific receptors on the islet β-cells, suppresses glucagon secretion, inhibits gastric emptying and reduces appetite [2]. Furthermore, chronic elevated concentrations of GLP-1 were shown to result in reduced hepatic fat accumulation and significantly lower TG concentrations in rat and mouse model [3]. However, after enzymatic degeneration via DPP4, which occurs within minutes following ingestion, only 10–20% of active GLP-1 remains in blood. Today, DPP4 inhibitors are employed in human medicine to prolong the beneficial incretin effects, in particular to improve insulin sensitivity, with the aim to treat type II diabetes [4].

In high-yielding dairy cows, the metabolic status around calving and the onset of lactation exhibits huge parallels to patients suffering from type II diabetes and non-alcoholic fatty liver diseases, as comprehensive physiological challenges are necessary to coordinate the metabolic alterations. In the transition from late pregnancy and early lactation, decreases in insulin concentration and peripheral insulin responsiveness suppress glucose consumption by peripheral, insulin-dependent tissues (skeletal muscle, adipose tissue) and thus enhance the availability of glucose for the insulin-independent mammary gland [5]. The adaption to the negative energy balance (NEB) is often related to metabolic dysfunctions, such as excessive lipid accumulation in the liver and ketosis [6–8], characterized by increased concentrations of non-esterified fatty acid (NEFA) and β-hydroxybutyrate (BHB). Due to the infiltration of fat, lesions in hepatic tissues appear and cause increased blood levels of specific enzymes, such as ω-glutamyl transferase (γ-GT), aspartate transaminase (AST) or glutamate dehydrogenase (GLDH) [9]. Furthermore, a fatty liver contributes the development of hepatic insulin resistance and influences body’s immune system negatively. In particular, the impacts of tumor necrosis factor-α and acute phase protein reactions are well studied in cows with fatty liver and its role in immune response [10,11].

Little is known about the metabolism of incretins and its interaction with DPP4 in ruminants. In contrast to monogastric species, the cow does not rely on glucose absorption in the small intestine but uses short chain fatty acids from ruminal fermentation for her energy supply with propionate as main substrate for gluconeogenesis. However, increasing dietary energy supply has been shown to enhance the secretion of GLP-1 in steers [12] and abomasal infusion of lipid and casein, but not glucose, increased the GLP-1 concentration in cows [13,14]. The fat-induced elevation in circulating GLP-1 is believed to play a role in the short-term control of feed intake in cattle [14,15], but the wide range of tissues expressing the GLP-1 receptor (gut segments, pancreas, spleen and kidney) suggest that GLP-1 may have multiple physiological functions beyond the control of feed intake [16]. The DPP4 expression and the circulating GLP-1 concentrations in blood depend on stage of lactation. While GLP-1 concentrations increase with onset of lactation, the expression of DPP4 decreases [13,17].

Taking the background information into consideration, DPP4 is a key enzyme in intermediary metabolism by regulating important glycemic pathways. Therefore, it was possible that DPP4 inhibitors could counteract typical ketotic processes in the dairy cow. Within the present research a DPP4 inhibitor (BI 14332) was employed to regulate typically increased parameters of bovine ketosis to the physiological range, respectively to compensate a distinct NEB. Therefore, we first established appropriate dosage of BI 14332 to effectively decrease DPP4 activity in plasma and liver from healthy lactating dairy cows. The second aim was to verify the efficacy of the derived dose and dosing regimen in cows with subclinical ketosis based on evaluation of...
various endpoints, such as clinical-chemical parameters and immune traits as well as liver lipid concentration.

Materials and Methods

Ethic statement

The experiments were approved by the competent authority, the lower Saxony state office for consumer protection and food safety (LAVES; Trial 1: file no. 33.9-42502-05-11A172, Trial 2: file no. 33.14-42502-04-11/0444; Oldenburg, Germany). The regulations of the German Animal Welfare Act (TierSchG) in its respective edition were met.

Experimental design

The investigations about the pharmacokinetics and pharmacodynamics (PK/PD) of BI 14332 were performed at the Clinic for Cattle, University of Veterinary Medicine in Hannover, Germany (Trial 1). The experiment aimed in evaluating the effectiveness of DPP4 inhibition in dairy cows with subclinical ketosis was carried out at the experimental station of the Institute of Animal Nutrition, Friedrich-Loeffer-Institute (FLI) in Braunschweig, Germany (Trial 2).

**Trial 1.** Six lactating and clinically healthy German Holstein cows (pluriparous) were treated with three different doses, i.e. 0.3, 1.0 and 3.0 mg/kg body weight (BW; injection volume: 0.01–0.1 mL/kg) of BI 14332 (n = 2/dosage; i.v.).

To evaluate the concentration of BI 14332 and the DPP4 activity in plasma, samples were collected 24 h before the injection, 0, 0.25, 0.5, 1, 2, 4, 6, 12, 24, and 48 h after the injection. In addition, liver samples were collected according to Starke et al. [18] 24 h before injection and 4, 24 and 48 h thereafter to evaluate the hepatic DPP4 activity.

For an in vitro activity assay, potassium EDTA plasma samples of three healthy Holstein Frisian cows were incubated with 0, 1, 3, 10, 30 and 100 nM BI 14332.

**Trial 2.** Using an animal model in which subclinical ketosis is induced [19]; the impact of DPP4 inhibition via BI 14332 on metabolic variables and on immune function was investigated. β-hydroxybutyrate concentrations between 1.2–2.5 mM in blood serum were defined as a subclinical ketotic status [20].

The chemical compositions of concentrate and total mixed ration are shown in Table 1. For more details regarding feeding management refer to Schulz et al. [19]. In brief, 20 pregnant and healthy German Holstein cows with a mean body condition score (BCS) of 3.16 ± 0.06 [21] were assigned to the experimental group. For the last six weeks of parturition, cows were fed with a high energetic ration (7.7 MJ NEL/kg dry matter). The allocation of cows with a BCS of at least 3.0 and a high energetic energy supply ante partum resulted in a higher condition at the time of calving (called “higher condition” cows; HC). The aim was to enhance lipomobilisation post partum. Immediately after calving, the energy supply by concentrate feeding was reduced first and raised stepwise (from 30% to 50% for the first three weeks of lactation).

Two cows from HC group were excluded from the experiment because of health problems, which were not due to the experimental design. Out of the 18 HC cows, 12 cows developed subclinical ketosis (serum BHB concentration ≥ 1.2 and < 2.5 mM). Six HC cows were treated with BI 14332 (HC-BI) over a period of 7 days (daily i.v. injections, 0.3 mg/kg BW/day) and the remaining six subclinical cows formed the control group (HC-Con) without treatment. Subclinical ketosis was diagnosed on day +3 (1 HC-Con cow), day +7 (5 HC-BI cows and 3 HC-Con cows) or on day +10 (1 HC-BI cow and 2 HC-Con cows), relative to calving. The remaining six HC cows were either affected with clinical ketosis (n = 3, BHB in serum > 2.5 mM) or stayed apparently healthy (BHB < 1.2 mM, n = 3). Blood samples were collected at day "0", i.e. 48 ± 7.2 days before expected calving, and then on day -14, -7, -3, +1, +3, +7,
+10, +14, +17, +21, +24, +28, +35, +42 and +56 (relative to calving) from the *V. jugularis*; liver biopsies were taken on day -14, +7, +21, +35 and +56.

Sample Preparation and Analysis

**Trial 1.** The DPP4 activity in plasma and liver was assessed by a semi-quantitative assay with fluorescence detection at the Institute for Clinical Research and Development (Mainz, Germany). The method was validated for the detection of DPP4 activity in human plasma samples [22]. The dose dependently inhibition of DPP4 by increasing BI 14332 in nanomolar concentrations (Fig 1) showed that DPP4 activity in bovine samples is detectable by the method as well. The fluorescence measured as relative fluorescence units (RFU) is equivalent to the DPP4 activity in the sample. The baseline/pre-dose activity was set to 100% and all other activities measured in blood samples of the individual cows were calculated as the respective percentage of the baseline DPP4 activity.

Homogenized liver samples (20 mg) were mixed with 450 μL DPP4 tissue buffer (25 mM HEPES, 140 mM NaCl, 80 mM MgCl₂ and 11.25 mM Triton x-100). The DPP4 tissue buffer containing liver material was collected in a vial, centrifuged (1780 g, 10 min, 4°C) and the resulting supernatant was stored at -80°C until analysis for DPP4 activity.

The concentration of BI 14332 in blood plasma was analyzed by the Pharmacelsus GmbH (Saarbrücken, Germany) via LC-MS/MS (Surveyor MS Plus HPLC system, Thermo Fisher Scientific), connected to a TSQ Quantum Discovery Max (Thermo Fisher Scientific) triple quad mass spectrometer. Data handling was done using the standard software Xcalibur 2.0.7.

| Table 1. Ingredients and chemical compositions of concentrate and total mixed ration. |
|----------------------------------|---------------------------------|-----------------|------------------|-----------------|
|                                  | Ante partum diet[^a^]           | Post partum diet[^b^] |
|                                  | Concentrate | TMR | Concentrate | TMR           |
| Ingredients, %                  |       |     |       |     |
| Wheat                            | 41.0  | 41.0 | 41.0  | 41.0 |
| Dried sugar beet pulp            | 30.5  | 30.3 | 30.3  | 30.3 |
| Rapeseed meal                    | 20.0  | 20.0 | 20.0  | 20.0 |
| Soybean meal                     | 6.5   | 6.5  | 6.5   | 6.5  |
| Vitamin/mineral premix           | 2.0[^c^] | 2.0[^d^] | 2.0[^c^] | 2.0[^d^] |
| Calcium carbonate                | -     | 0.2  | -     | 0.2  |
| Dry matter (DM), g/kg            | 877   | 489  | 875   | 393  |
| Nutrients [g/kg DM]              |       |      |       |      |
| Crude ash                        | 58    | 55   | 62    | 56   |
| Crude protein                    | 197   | 140  | 202   | 122  |
| Ether extract                    | 27    | 33   | 28    | 32   |
| Crude fibre                      | 101   | 163  | 72    | 194  |
| Acid detergent fibre (ADF)       | 136   | 199  | 96    | 222  |
| Neutral detergent fibre (NDF)    | 279   | 394  | 222   | 431  |
| Energy[^e^], MJ NEL/kg DM        | 8.6   | 7.7  | 8.7   | 7.0  |

[^a^]Total mixed ration (TMR) on dry matter (DM) basis (40% roughage (75% corn silage, 25% grass silage) + 60% concentrate.
[^b^]TMR on DM basis (70% roughage (75% corn silage, 25% grass silage) + 30% concentrate.
[^c^]Per kg of mineral feed: 10g Ca, 60g P, 120g Na, 60g Mg, 800,000 IU vitamin A, 100,000 IU vitamin D₃, 2500mg vitamin E, 4000mg Mn, 6000 mg Zn, 1250mg Cu, 100mg I, 35mg Co, 50mg Se
[^d^]Per kg of mineral feed: 170g Ca, 50g P, 120g Na, 45g Mg, 800,000 IU vitamin A, 100,000 IU vitamin D₃, 4000mg vitamin E, 4000mg Mn, 6000mg Zn, 1300mg Cu, 120mg I, 35mg Co, 40mg Se
[^e^]Calculation based on nutrient digestibilities measured with wethers (GIE, 1991) and values from feed tables (DLG, 1997)

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**Trial 2.** Clinical chemistry [BHB, NEFA, triglycerides (TG), glucose, γ-glutamyl transferase (γ-GT), aspartate transaminase (AST), glutamate dehydrogenase (GLDH)] was assessed in serum using photometric methods (Eurolyser, Type VET CCA, Eurolyser Diagnostica GmbH). A radioimmunoassay was used to quantify the serum insulin concentrations (IM3210, Insulin IRMA KIT, Immunotech, Beckman Coulter). This immunoradiometric test was a "sandwich" type assay. The antibodies used (mouse monoclonal) were directed against two different insulin epitopes. The assay was performed according to the manufacturer’s instructions. The intra-assay CV was 7.6%, and the inter-assay CV was 10.7%. The lowest detection limit was 3.95 μU/ml. Haptoglobin (Hp) was measured by ELISA as described elsewhere [23] and the total lipid content in liver samples (~100 mg) was assessed using a gravimetric method [18].

The daily dry matter intake (DMI) was recorded for the whole experimental time (computerized feeding station: Type RIC, Insentec). Milking took place twice a day at 05.30 and 15.30. Milk yield was recorded using automatic milk counters (Lemmer Fullwood GmbH).

Hematological analyses were performed in EDTA whole blood using an automatic analyzer (Celltac α MEK-6450, Nihon Kohden, Qinlab Diagnostik).

Functional tests (ex vivo) of peripheral blood mononuclear cells (PBMC) were performed in samples from days -14, +7, +10, +14, +21 and +56 (relative to calving) using the Alamar Blue assay (AB). Concanavalin A (ConA, 2.5 μg/mL final, Sigma-Aldrich) was used as mitogen to stimulate T-lymphocytes. Further details were described previously [24].

For the calculation of CD4+/CD8+ T-cell population and its CD4+/CD8+ ratio from data generated by flow cytometry, days relative to calving where pooled in accordance to treatment [day “0” and day -14 (“ante-partum”), two days of treatment (“treatment”), after treatment, i.e.
day +17 until day +28 post-partum ("2 weeks post treatment") and day +35, +42 and +56 post-partum ("end of trial"). A detailed description of the measurements is provided by Stelter et al. [25]. Samples were double stained with monoclonal antibodies for CD4⁺ (mouse anti bovine CD4:FITC) and CD8⁺ (mouse anti bovine CD8:RPE) or the corresponding isotype controls (mouse IgG2a negative control: RPE and mouse IgG2b negative control: FITC; all AbD Serotec).

Statistics and Calculations

**Trial 1.** The pharmacokinetic parameters were performed using non-linear regression via STATISTICA 10 [26]. The time course of plasma concentration of BI 14332, $C_p$, was expressed by a sum of two exponential functions:

$$C_p = \sum_{i=1}^{n} a_i e^{-b_i t}$$

where $a_i$ and $b_i$ are hybrid coefficients and exponential terms, $t$ is time, and $n$ is the number of exponential terms. From the data obtained, area under the concentration-time curves from 0 to 24 h (AUC), terminal half-life ($t_{1/2}$), total body clearance from 0 to 24 h ($Cl_{24h}$), the Volume of distribution ($V_d$), and the average steady state concentration ($Css$) were calculated.

Area under the curve of DPP4 activity in plasma and liver from 0 to 24 h was calculated using the linear trapezoidal rule:

$$AUC = \sum_{n=1}^{N} \frac{C_n + C_{n+1}}{2} \left( t_{n+1} - t_n \right)$$

**Trial 2.** Insulin sensitivity was estimated by the Revised Quantitative Insulin Sensitivity Check Index (RQUICKI) [27]:

$$RQUICKI = \frac{1}{\log \text{Insulin}[\mu U/mL] + \log \text{Glucose}[mg/dL] + \log \text{NEFA}[mmol/L]}$$

The results of the ex vivo examinations of PBMC were expressed as stimulation index (SI), defined as ratio between the fluorescence in the AB assay of ConA stimulated and nonstimulated PBMC:

$$SI = \frac{\text{Fluorescence of ConA stimulated PBMC}}{\text{Fluorescence of nonstimulated PBMC}}$$

For statistical analyses the SAS software package [28] was used. Evaluation of goodness of fit was carried out using the corrected Akaike information criterion. All parameters evaluated were compared as dependent variable by the MIXED procedure with a compound symmetry covariance structure. Treatment (HC-Con vs. HC-BI) was considered as fixed factor and sampling dates (time) as a repeated effect, and their respective interaction were included into the model. All results are presented as least square means (LSmeans) and standard errors (SE). Effects were declared to be significant when $P$-values were $< 0.05$ after Tukey test for post-hoc analysis, whereas a trend was noted when $0.05 < P < 0.10$.

Data evaluation of hematology, proliferative capability (SI) and clinical chemistry based on pooled sampling days ("week of treatment/observation", "1st week after treatment/observation" and "2nd week after treatment/observation"). The day with first occurrence of serum BHB concentration $\geq 1.2$ mM (day of classification) was set as covariate. Data evaluation of milk
yield and DMI based on weekly mean values. The first week of lactation was set as covariate. For the proliferative capability, the SI of day +7 post partum was set as covariate. The remaining variables (liver lipid content, parameters of glycemic control, phenotyping T-lymphocytes) were analyzed in accordance to the evaluated sampling days, as described above.

Results

Trial 1

Investigations in vitro. The DPP4 activity in EDTA plasma samples decreased significantly starting at a concentration of 3 nM (± 1.407 ng/mL) BI 14332 as shown in Fig 1. At 100 nM (± 46.9 ng/mL) the remaining DPP4 activity was 15.1%.

Pharmacokinetics and pharmacodynamics. The single administration of BI 14332 at 0.3, 1.0 or 3.0 mg/kg BW was well tolerated and a clear BI 14332 plasma concentration-dependent inhibition of the DPP4 activity both in plasma and liver was noted (Fig 2). The PK/PD variables of BI 14332 and DPP4 activity in plasma and liver are represented in Table 2. The AUC regarding BI 14332 in plasma were dose-dependently increased. The $t_{1/2}$ of BI 14332 was

![Fig 2. Inhibition of plasma and liver dipeptidyl peptidase-4 (DPP4) activities after injection of BI 14332. BI 14332 was administered in a single dose of 3 [square], 1 [triangle] and 0.3 [circle] mg/kg body weight in dairy cows (n = 2/group). Plasma samples (V. jugularis) were taken 24 h before and immediately before (time zero “0”) injection, as well as 0.25, 0.5, 1, 2, 4, 6, 12, 24 and 48 h post injection (upper shape). Liver was biopsied 24 h before injection, as well as 4, 24 and 48 h thereafter (lower shape). doi:10.1371/journal.pone.0136078.g002]
highest for the 0.3 mg/kg BW dosage group, with a 10.5 to 23 h range. The \( V_d \) and \( Cl_{24h} \) were greatest when 3 mg/kg BW was applied. The \( C_{ss} \) decreased dose-dependently, starting with the highest dosage of BI 14332. Plasma DPP4 activity (Fig 2A) was significantly inhibited by BI 14332 at all dosages with a remaining maximum activity of 14%, which was in line with an inhibitory power of 86% (1 mg/kg BW; 15 min after injection). The single dose of 1 mg/kg also showed the lowest inhibition at 24 h after injection (~ 70%), while inhibition by the other two dose groups were greater and quite similar (81–87%). Forty-eight hours after injection, the cows treated with 0.3 and 3 mg/kg still had an inhibition of DPP4 activity of about 82% versus 74% for the 1 mg/kg dosage group (\( P < 0.05 \)). In liver, the AUC of DPP4 activity was decreased with increasing dosage (Fig 2B).

The relationship between the BI 14332 concentration in plasma and the corresponding DPP4 activity in plasma and liver is depicted in Fig 3, well approximated by a power function. With increasing plasma concentrations of BI 14332 (x-axis), a negative slope for DPP4 activity (y-axes) was observed in liver: \( y = 7.72x^{0.589} \) (\( r^2 = 0.72 \)) and in plasma: \( y = 935.31x^{0.081} \) (\( r^2 = 0.76 \)).

Trial 2

Clinical chemistry and hepatic lipid content. An overview of the clinical chemical parameters is given in Table 3. A significant group \( \times \) time interaction was shown for NEFA, TG

**Table 2. Pharmacokinetic parameters of BI 14332 und Dipeptidy peptidase-4 (DPP4) in plasma and liver of six healthy German Holstein cows treated with different dosages of BI 14332 [3, 1 and 0.3 mg/kg body weight (BW); n = 2/dosage group].**

| BI 14332 | Dosage | 3 mg/kg BW | 1 mg/kg BW | 0.3 mg/kg BW |
| --- | --- | --- | --- | --- |
| | Cow 1 | Cow 2 | Mean | Cow 3 | Cow 4 | Mean | Cow 5 | Cow 6 | Mean |
| **Plasma** | | | | | | | | | |
| AUC\(_{24h}\) [ng\( \cdot \)h/mL] | 2076 | 2784 | 2430 | 1769 | 773 | 1271 | 809 | 783 | 796 |
| \( t_{1/2} \) [min] | 198 | 289 | 243 | 116 | 173 | 144 | 630 | 1386 | 1008 |
| \( V_d \) [L/kg BW] | 17.07 | 30.13 | 23.60 | 5.21 | 12.73 | 8.97 | 13.37 | 20.56 | 16.97 |
| Cl\(_{24h}\) [mL/kg/min] | 24.09 | 17.96 | 21.02 | 9.42 | 21.55 | 15.49 | 6.18 | 6.39 | 6.28 |
| \( C_{ss} \) [ng/mL] | 86.49 | 115.99 | 101.24 | 71.94 | 30.70 | 51.32 | 33.69 | 32.63 | 33.16 |
| **DPP4** | | | | | | | | | |
| AUC\(_{24h}\) [RFU/h] | 20491 | 16560 | 18525 | 21034 | 20135 | 20584 | 15729 | 17914 | 16822 |
| \( \Delta_{15min} \) [%] | 90 | 91 | 91 | 86 | 86 | 86 | 88 | 90 | 89 |
| \( \Delta_{24h} \) [%] | 81 | 85 | 83 | 66 | 73 | 70 | 84 | 87 | 85 |
| **Liver** | | | | | | | | | |
| AUC\(_{24h}\) [\( \mu \)g\( \cdot \)h/mL] | 41 | 74 | 43 | 90 | 45 | 68 | 63 | 112 | 101 |
| \( \Delta_{4h} \) [%] | 96 | 94 | 94 | 89 | 89 | 89 | 77 | 79 | 78 |
| \( \Delta_{24h} \) [%] | 65 | 71 | 68 | 40 | 44 | 42 | 35 | 31 | 33 |

\( a \)Pharmacokinetic parameters of BI 14332 were evaluated via bi-exponential function [26]; AUC for DPP4 activity in plasma and liver was calculated using the trapezoidal rule.

\( b \)BI 14332 was administrated intravenously (i.v.); plasma samples were taken 24 h before injection, 0, 0.25, 0.5, 1, 2, 4, 6, 12, 24 and 48 h after injection; liver samples were taken 24 h before injection and 4, 24 and 48 h thereafter.

\( c \)DPP4 activity in liver was normalized to the total protein content of the samples. AUC, area under the curve from 0 to 24 h; \( t_{1/2} \), terminal half-life; \( V_d \), Volume of distribution; Cl\(_{24h}\), Clearance from 0 to 24 h; \( C_{ss} \), average steady state concentration; \( \Delta_{15min}/\Delta_{4h}/\Delta_{24h} \): Inhibitory power of BI 14332 regarding DPP4 activity, calculated as difference before BI 14332 application and the first sample post injection (i.e. 15 min post injection in plasma and 4 h post injection in liver) and 24 h after injection, respectively.

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and GLDH. All three variables changed significantly with time and were additionally influenced by treatment. For TG, this was indicated by a significantly lower concentration in the HC-BI cows during the 1st week after treatment compared to the HC-Con cows. For NEFA, there was a significant decrease in the concentration between the week of treatment to the 1st and the 2nd week post treatment, only within the HC-BI group, while GLDH increased significantly in the HC-Con cows and peaked in the 2nd week post observation. Further time-dependent alterations were detected for glucose, insulin, AST and γ-GT. Glucose and insulin concentrations were significantly greater two weeks post observation than during observation in the HC-Con group. For more detailed evaluation of variables describing glycemic control, Fig 4 shows RQUICKI and the variables necessary to calculate the index (NEFA, glucose, insulin). The figure reveals significant changes relative to calving and in accordance to treatment. On day +10 post partum, NEFA, glucose and insulin peaked within the HC-BI group, while RQUICKI decreased to a nadir at that day. For the HC-Con cows there was the opposite effect as substantiated by the significant interaction between group and day for RQUICKI. Starting on day +14 until day +21 post partum, RQUICKI differed markedly between the experimental groups, with higher values for the HC-BI cows.

The BHB concentrations were not different between the groups and the same was true for Hp. The greatest Hp concentrations were detected at the day of classification in both groups; thereafter the concentration decreased continuously. Albeit not reaching the level of significance, the decrease of the Hp concentration seemed to be faster in HC-BI than in the HC-Con cows.

There was also no significant difference for the total liver lipid content between groups (data not shown), even though HC-Con cows had slightly higher lipid contents than the HC-BI cows at all days evaluated post partum (Δ = 13.8 mg/g). For both groups the total hepatic lipid content differed significantly with time and highest contents were detected on day +7 after calving with 161 mg/g (HC-BI) and 175 mg/g (HC-Con).
Table 3. Effects of dipeptidyl peptidase-4 (DPP4) inhibition via BI 14332 to blood serum variables of clinical chemistry and insulin sensitivity of cows with subclinical ketosis (LSmeans ±SE).

| Parameter          | HC-BI (n = 6) | HC-Con (n = 6) | Probability |
|--------------------|---------------|----------------|-------------|
|                    | Day of        | 1st week after | 2nd week after | Day of       | 1st week after | 2nd week after |
|                    | classification | treatment      | treatment    | classification| observation    | observation    |
| BHB [mM]           | 1.63 ± 0.65   | 1.77 ± 0.38    | 1.56 ± 0.31  | 1.51 ± 0.23  | 1.44 ± 0.31    | 1.15 ± 0.30    | 1.12 ± 0.32    | 0.509         | 0.166         | 0.520         |
| NEFA [mM]          | 0.92 ± 0.28   | 1.09 ± 0.10    | 0.56 ± 0.08  | 0.94 ± 0.23  | 0.94 ± 0.08    | 0.75 ± 0.08    | 0.55 ± 0.08    | 0.820         | <             | 0.026         |
| Triglyceride [mg/dL]| 11.07 ± 2.40  | 11.20 ± 1.10   | 9.03 ± 0.74  | 9.57 ± 0.85  | 12.34 ± 3.69   | 11.37 ± 0.87   | 14.70 ± 0.81   | 11.02 ± 0.88  | 0.014         | 0.123         | 0.002         |
| Glucose [mg/dL]    | 50.68 ± 9.22  | 55.09 ± 4.49   | 57.64 ± 3.72 | 59.86 ± 3.97 | 63.05 ± 8.52   | 51.70 ± 3.93   | 60.84 ± 3.74   | 62.58 ± 3.89  | 0.876         | 0.018         | 0.404         |
| Insulin [mU/mL]    | 5.20 ± 2.72   | 4.58 ± 1.79    | 5.78 ± 1.29  | 8.63 ± 1.44  | 6.70 ± 5.62    | 5.49 ± 1.46    | 7.17 ± 1.43    | 11.21 ± 1.48  | 0.313         | 0.001         | 0.802         |
| RQUICKI            | 0.44 ± 0.04   | 0.43 ± 0.02    | 0.47 ± 0.01  | 0.44 ± 0.02  | 0.41 ± 0.04    | 0.43 ± 0.02    | 0.42 ± 0.02    | 0.39 ± 0.02   | 0.129         | 0.127         | 0.160         |
| Haptoglobin [mg/mL]| 1.56 ± 1.63   | 0.31 ± 0.30    | 0.13 ± 0.21  | 0.12 ± 0.23  | 1.27 ± 1.22    | 0.76 ± 0.24    | 0.53 ± 0.23    | 0.32 ± 0.24   | 0.168         | 0.385         | 0.817         |
| AST [IU/l]         | 108.03 ± 35.3 | 117.00 ± 10.8  | 104.86 ± 8.18| 86.43 ± 8.92 | 96.79 ± 21.0   | 112.65 ± 9.07  | 105.11 ± 8.68  | 92.83 ± 9.19  | 0.941         | 0.005         | 0.772         |
| γ-GT [IU/l]        | 18.49 ± 3.41  | 21.67 ± 9.01   | 31.50 ± 8.15 | 40.30 ± 8.37 | 19.95 ± 4.18   | 22.94 ± 8.44   | 28.54 ± 8.30   | 38.85 ± 8.47  | 0.928         | 0.001         | 0.868         |
| GLDH [IU/l]        | 8.13 ± 1.56   | 19.89 ± 7.72   | 29.75 ± 6.56 | 27.08 ± 6.88 | 9.42 ± 2.59    | 13.59 ± 6.90   | 20.92 ± 6.67   | 39.41 ± 6.97  | 0.918         | 0.003         | 0.028         |

With first occurrence of serum β-hydroxybutyrate (BHB) concentration ≥ 1.2 mM cows were treated with BI 14332 (HC-BI) or stayed untreated as control group (HC-Con). BI 14332 was applied once a day over a period of 7 days (i.e., 0.3 mg/kg body weight). Subclinical ketosis was diagnosed on day +3 (1 cow), day +7 (8 cows) or day +10 (3 cows) after calving. The day of classification (mean ± SD), which was the day with first occurrence of BHB values ≥ 1.2 mM were set as covariate, integrated in the MIXED procedure of SAS [24] with group and time as fixed factors (P ≤ 0.05; Tukey test). Significant values are shown in bold.

NEFA, non-esterified fatty acids; RQUICKI, revised quick insulin sensitivity index; AST, aspartate aminotransferase; γ-GT, γ-glutamyltransferase; GLDH, glutamate dehydrogenase
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Fig 4. Concentrations of non-esterified fatty acids (NEFA), glucose and insulin in serum, and insulin sensitivity (RQUICKI) in cows with subclinical ketosis. With the first occurrence of serum β-hydroxybutyrate concentrations ≥ 1.2 mM, cows were treated with BI 14332 [(—) n = 6] or stayed untreated as control [(--) n = 6]. Within the BI 14332 treatment group subclinical ketosis was diagnosed on day +7 (5 cows) and on day +10 (1 cows), relative to calving. Dosage of BI 14332 was 0.3 mg/kg body weight, applied i.v. once a day over a period of 7 days. The statistical analysis included group (BI 14332 treatment vs. control), experimental day (1st day post partum until 56th day post partum), and the interaction (P < 0.05, Tukey test). Experimental day differed significantly for all parameters. [NEFA (diamond), Glucose (triangle), Insulin (circle), RQUICKI (square)].

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DMI and milk yield. Statistical evaluation of dry matter intake and milk yield during lactation is shown in Table 4. Data offered a significant time effect for the variables. Milk yield and DMI increased continuously for both subgroups. A group effect, respectively a significant group*time interaction, was not determined for the named parameters.

Hematology and cell function. Variables of the hematology and the SI of PBMC are shown in Table 5. White blood cells (WBC) were significantly influenced by time and time*group. During treatment, leucocytes and granulocytes counts were significantly higher in the HC-BI than in the HC-Con group. The decrease of the named parameters from treatment to the 1st and 2nd week after treatment was significant in the HC-BI group. The proliferative capability of PBMC in the ex vivo assay did not change over time. T-cell-phenotyping revealed that the CD4+/CD8+ ratio of subclinically cows tended to be higher for the HC-Con versus the HC-BI group (P = 0.059, Table 6). For the HC-Con cows the increase from ante partum to treatment period was significant; due to a selective increase of the CD4+ T-cell population (~ 11%) and a slight decrease of the CD8+ T-cell population (~ 10%). In the HC-BI group, both subpopulations decreased (CD4+: ~ 16%, CD8+: ~ 26%) during treatment. Both experimental groups showed significant time effect with highest ratios during observation (HC-Con: 3.28) and two weeks after treatment (HC-BI: 2.60). The individual CD4+ and CD8+ cell populations differed significantly over the time, but neither group nor the group*time interaction were significant.

Discussion

Most of the findings about inhibition of DPP4 are related to type II diabetes in human patients, opening up new perspectives in therapy. The ketogenic metabolic status of high yielding dairy cows during early lactation is characterized by some similarities with the metabolic situation of type II diabetes and fatty liver in humans [29].

The first aims of the current investigation were (1) to show that BI 14332 is an effective DPP4 inhibitor and (2) to find the optimal dosage of BI 14332. Therefore, all doses resulted in an inhibition of DPP4 activity in plasma and liver without any indication for adverse side-effects. According to the PK/PD results, we considered the dosage of 0.3 mg/kg BW as most suitable. Albeit the dosage of 3 mg/kg BW yielded the highest AUC of BI 14332, the plasma AUC of DPP4 activity was quite similar in the highest and the lowest dosage group. The
Table 5. Effects of dipeptidyl peptidase-4 (DPP4) inhibition via BI 14332 to hematological variables and the proliferative capability of PBMC (LSmeans ± SE).

| Parameter       | HC-BI (n = 6) | HC-Con (n = 6) | Probability     |
|-----------------|---------------|----------------|-----------------|
|                 | Day of        | 1<sup>st</sup> week | 2<sup>nd</sup> week | Day of        | 1<sup>st</sup> week | 2<sup>nd</sup> week | group | time | Group x time |
|                 | Classification | after treatment | after treatment | Classification | after observation | after observation |       |      |             |
| SI              | 6.69 ± 1.29   | 6.05 ± 0.79    | 6.26 ± 0.56     | 3.69 ± 1.42    | 7.27 ± 1.62     | 6.73 ± 0.75      | 6.49 ± 0.64     | 6.80 ± 1.37 | 0.138 0.591 0.453 |
| Leucocytes [10<sup>9</sup>/μL] | 9.75 ± 2.79   | 10.17 ± 0.65   | 7.59 ± 0.47     | 7.14 ± 0.51    | 7.35 ± 3.07     | 7.33 ± 0.52      | 7.48 ± 0.50     | 7.05 ± 0.56  | 0.105 0.005 0.009 |
| LY [10<sup>9</sup>/μL] | 3.30 ± 1.07   | 2.79 ± 0.22    | 2.86 ± 0.19     | 2.89 ± 0.20    | 2.93 ± 0.41     | 2.95 ± 0.20      | 2.97 ± 0.19     | 3.09 ± 0.20  | 0.549 0.642 0.927 |
| GR [10<sup>9</sup>/μL] | 5.87 ± 2.01   | 6.74 ± 0.66    | 4.01 ± 0.48     | 3.75 ± 0.52    | 3.77 ± 2.95     | 4.00 ± 0.53      | 3.99 ± 0.51     | 3.37 ± 0.57  | 0.106 0.002 0.015 |
| EO [10<sup>9</sup>/μL] | 0.40 ± 0.24   | 0.33 ± 0.12    | 0.39 ± 0.09     | 0.29 ± 0.10    | 0.45 ± 0.26     | 0.27 ± 0.10      | 0.34 ± 0.10     | 0.40 ± 0.11  | 0.987 0.730 0.426 |
| Erythrocytes [10<sup>12</sup>/μL] | 5.92 ± 0.42   | 5.81 ± 0.21    | 5.69 ± 0.19     | 5.53 ± 0.20    | 6.07 ± 0.82     | 5.90 ± 0.20      | 5.94 ± 0.19     | 5.87 ± 0.20  | 0.407 0.238 0.468 |
| HGB [g/dL]      | 10.82 ± 0.68  | 10.23 ± 0.42   | 9.90 ± 0.37     | 9.69 ± 0.39    | 10.52 ± 1.18    | 10.40 ± 0.39     | 10.38 ± 0.39    | 10.31 ± 0.40  | 0.435 0.331 0.551 |
| HCT [%]         | 36.30 ± 2.61  | 34.60 ± 1.36   | 33.48 ± 1.20    | 32.27 ± 1.24   | 35.07 ± 4.09    | 34.70 ± 1.26     | 34.59 ± 1.24    | 34.43 ± 1.30  | 0.510 0.186 0.347 |
| Platelets [10<sup>9</sup>/μL] | 514 ± 127     | 117 ± 653      | 1029 ± 515      | 1631 ± 550     | 453 ± 48        | 835 ± 561        | 862 ± 538      | 909 ± 583    | 0.933 0.211 0.276 |

With first occurrence of serum β-hydroxybutyrate ≥ 1.2 mM cows were treated with BI 14332 (HC-BI) or stayed untreated as control (HC-Con). BI 14332 was applied once a day over a period of 7 days (intravenous, 0.3 mg/kg body weight). Subclinical ketosis was diagnosed on day +3 (1 cow), day +7 (8 cows) or on day +10 (3 cows) after calving.

The day of classification (mean ± SD), which was the day with first occurrence of BHB values ≥ 1.2 mM were set as covariate, integrated in the MIXED procedure of SAS [24] with group and time as fixed factors (P ≤ 0.05; Tukey test). Significant values are shown in bold.

SI, stimulation index (ratio between the fluorescence in the Alamar Blue assay of concanavalin A-stimulated and unstimulated PBMC); LY, lymphocytes; GR, granulocytes; EO, eosinophile granulocytes; HGB, hemoglobin; HCT, hematocrit.

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terminal t1/2 is the time required to halve the plasma concentration after reaching a steady state equilibrium and has to be known to provide an appropriate length for the dosing interval [30], which was sought to be 24 h. Results confirm dosing decision of 0.3 mg/kg BW, as t1/2 and Cl24h, i.e. the ability to eliminate a drug [31], offered the longest retention period in blood.

The aim of trial 2 was to test whether inhibiting DPP4 may affect blood variables associated with lipid metabolism and glycemic control under catabolic condition. The results showed that the impact of DPP4 inhibition was less in cows of subclinical ketosis, respectively during early lactation. Significant changes and positive impacts were limited to the time after treatment (Table 3; NEFA, TG, GLDH). Lower TG concentrations were also observed by Ben-Shlomo et al. [3]. The researches show significant reduced levels in DPP4-deficient rats and explain a GLP-1 mechanism in liver which induces a signal for a low energy state. Therefore, protein kinase activities involved in the pathway effect a reduced expression of lipogenesis-related genes. Effects of DMI and milk yield showed that there was no difference between the sub-groups (Table 4), which could have explained a lower lipolysis for the HC-BI cows. Therefore, the decrease in NEFA was not caused by a higher DMI or a lower milk yield for those cows compared to the HC-Con cows.

RQUICKI, as surrogate marker to assess insulin sensitivity, includes NEFA in the equation and the lower NEFA concentrations in HC-BI vs. HC-Con rather than the ones of glucose and insulin affected RQUICKI. However, differences between the groups were limited to diverging reaction over time, i.e. the interaction of time and treatment. At day +14 post partum RQUICKI was greater in the BI 14332 treated animals. At this stage, approximately during the middle of the treatment, a steady state plateau is reached and therapeutic efficacy is assumingly complete [32]. In a further study, RQUICKI remained unaffected during subclinical ketosis [19]. We conclude, that RQUICKI may be not sensitive enough to investigate alterations of insulin sensitivity in case of subclinical ketosis, but it is more likely that insulin sensitivity is not, or even very less, influenced by subclinical ketosis and a diminished sensitivity is limited to physiological changes around calving. Nevertheless RQUICKI was sensitive enough to show a transient improvement of insulin sensitivity by BI 14332. Taking into account that DPP4 inhibition was significant (Fig 2, Table 2) and with respect to investigations made in humans and rodents, one could assume that the retention period of active GLP-1 in the periphery was prolonged. If that is also true in cows, it may explain the positive impacts on lipaemic control. However, the support was not strong enough to affect the entire metabolism.

The fact that effects of DPP4 inhibition were marginal may be linked to the short duration of the treatment period [2]; which was possibly reflected by GLDH. Changes in GLDH became evident only late [19,33] and indicated less hepatic lesions in the second week after BI 14332
treatment. This was not evidenced by a reduced level of hepatic fat in the HC-BI cows. A prolonged treatment period might affect hepatic infiltrations of lipids in a stronger way.

Another aim of trial 2 was to investigate if the inhibition of DPP4 via BI 14332 has immune-modulating effects. The prescribing information of sitagliptin, the first DPP4 inhibitor for clinical use, reports a slight increase in WBC, primarily due to a small increase of neutrophil granulocytes counts [34]. The HC-BI cows had already higher WBC values before the onset of the treatment, with \( \Delta = 2.40 \cdot 10^3/\mu l \) at the day of classification compared to cows of control group. Therefore, it was questionable, if significant changes of WBC and GR counts were indeed related to BI 14332 or just coincided with time-related alterations around calving. Our investigations regarding the proliferative capability of PBMC showed similar results and immune-modulatory alterations were not evident.

Studies employing DPP4 inhibitors showed that T-cell proliferation and cytokine production is inhibited by impaired DNA synthesis [35]. In contrast, Anz et al. [36] showed results, similar to the present. None of the tested DPP4 inhibitors impaired key parameters of the innate and adaptive immune response, which were included in the present study to assess drug safety.

For the current investigation, immune-modulatory effects were limited to differences in CD4+/CD8+ ratio, which tended to be higher for the HC-Con cows (vs. HC-BI cows), with a significant increase after calving (~ 30%) up to 3.28. A ratio up to 2.5 suggests a physiological situation, while an increased ratio may indicate an immune dysregulation [37,38]. Furthermore it is known that DPP4 is expressed predominantly on T-lymphocytes and most of the T-cells expressing DPP4 belong to the CD4+ population (~ 56%) [39]. The inhibition of DPP4 activity may lower the expression of CD4+ and CD8+ T-cells. This was more pronounced for CD8+ cells and led to a better CD4+/CD8+ ratio. It suggests an impaired immune defense after calving, when the need for an appropriate defense is highest. In view of the concentrations of the acute-phase protein Hp that is commonly used as marker of inflammation, a beneficial effect of BI 14332 on the immune defense was not supported. Haptoglobin is assumed to be elevated by tissue lesions occurring during birth and by the general proinflammatory situation for the time around calving [40]. It positively correlates with BHB, NEFA and TG [10,24,41]. The minor impact of the DPP4 inhibition regarding these variables may explain for the insignificant differences in Hp concentrations between groups. Nevertheless, in numerical terms alone, within the HC-BI group Hp dropped markedly by about 80% (vs. ~ 40% for HC-Con cows; Table 3) during the treatment. Thus a prolonged treatment period together with a concomitant improvement of hepatic lipid metabolism may also reduce Hp.

Conclusions

The DPP4 activity was determined in plasma and liver samples of dairy cows. The DPP4 inhibitor BI 14332 reduced the enzymatic activity in vivo and showed a fast onset and a long lasting inhibition of DPP4. However, the DPP4 inhibition did not improve the metabolic disarrangements related to subclinical ketosis. Albeit an improved lipaemic control was observed, as NEFA and TG were decreased after treatment. Unfortunately, the support was not strong enough to affect main markers of ketosis (BHB, hepatic lipid content).

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Author Contributions
Conceived and designed the experiments: SD UM DR JR GB. Performed the experiments: KS JF MP MM HS. Analyzed the data: KS JF DR. Wrote the paper: KS SD HS.

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