Chronic Cannabidiol Administration Fails to Diminish Blood Pressure in Rats with Primary and Secondary Hypertension Despite Its Effects on Cardiac and Plasma Endocannabinoid System, Oxidative Stress and Lipid Metabolism

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Received: 4 February 2020; Accepted: 13 February 2020; Published: 14 February 2020

Abstract: We investigated the influence of cannabidiol (CBD) on blood pressure (BP) and heart rate (HR) in spontaneously (SHR) and deoxycorticosterone (DOCA-salt) hypertensive rats. Hypertension was connected with increases in cardiac and plasma markers of lipid peroxidation in both models, whereas cardiac endocannabinoid levels decreased in SHR and increased in DOCA-salt. CBD (10 mg/kg once a day for 2 weeks) did not modify BP and HR in hypertension but counteracted pro-oxidant effects. Moreover, it decreased cardiac or plasma levels of anandamide, 2-arachidonoylglycerol and oleoyl ethanolamide in DOCA-salt and inhibited the activity of fatty acid amide hydrolase (FAAH) in both models. In the respective normotensive control rats, CBD increased lipid peroxidation, free fatty acid levels and FAAH activity. In conclusion, chronic CBD administration does not possess antihypertensive activity in a model of primary and secondary (DOCA-salt) hypertension, despite its antioxidant effect. The latter may be direct rather than based on the endocannabinoid system. The unexpected CBD-related increase in lipid peroxidation in normotensive controls may lead to untoward effects; thus, caution should be kept if CBD is used therapeutically.

Keywords: 2-arachidonoylglycerol; anandamide; cannabidiol; cannabinoid receptor; SHR; DOCA-salt; endocannabinoids; oxidative stress

1. Introduction

Cannabidiol (CBD) is one of the most abundant cannabinoids derived from the Cannabis sativa plant and devoid of a psychoactive effect [1,2]. CBD binds to cannabinoid CB₁ and CB₂ receptors with much lower affinity than ∆⁹-tetrahydrocannabinol (THC) [3] and interacts with GPR18, GPR55 and TRPV1 receptors [4]; it possesses a very marked antioxidant effect [5–7]. CBD is licensed for the treatment of some types of childhood epilepsy (Dravet and Lennox-Gastaut syndrome) in the United States.
States \[4,8\] and, in combination with THC, for the treatment of multiple sclerosis-associated spasticity in Canada and in the European Union \[4\]. In addition, a potential therapeutic action of CBD is being considered in anxiety disorders, schizophrenia, depression, Alzheimer’s disease, Parkinson’s disease, pain, cancer, inflammatory and autoimmune diseases and diabetic complications \[2,4,9\].

CBD may become a strategy also for the treatment of cardiovascular diseases, including hypertension \[3,9\]. To date, blood pressure-lowering effects of CBD were observed under stress conditions in humans \[10–12\] and in stressed animals \[13,14\]. However, the effect of CBD on the blood pressure of hypertensive individuals has been studied in one study only; in a paper on conscious spontaneously hypertensive rats \[15\], a single intraperitoneal dose of CBD (10 mg/kg) failed to affect blood pressure.

Hypertension is a disease with a complex pathomechanism, which includes, among others, changes in the endothelium and redox balance, both within the heart and blood vessels \[16,17\]. CBD is suggested to be a potential positive modulator of hypertension thanks to its vasodilatory action \[3,9,11,18\]. Another property that may be of key importance in a potential antihypertensive activity of CBD is its impact on oxidative stress. Attenuation of oxidation and/or nitration parameters by CBD was observed in acute experiments on human endothelium cells treated with high glucose \[19\], on the liver of mice subjected to ischemia/reperfusion \[20\] and on mouse hippocampal cells subjected to oxygen plus glucose deprivation/reperfusion \[21\]. Similar beneficial effects were also obtained in chronic experiments on the heart \[22\] and retina \[23\] from diabetic mice, on mouse hepatic cells with ethanol-induced liver injury \[24,25\], on the heart from doxorubicin-treated mice \[26\] and rats \[27\] and on the heart and other tissues of rats with sepsis \[28\].

The mechanism of CBD in the latter studies is complex and probably results from direct antioxidant properties \[3,29\] but may also be related to an effect on the endocannabinoid system, which is important for the modulation of oxidative stress \[30,31\]. CB1 receptors are mainly associated with its promotion \[32–34\], whereas CB2 \[35–39\] and GPR18 \[40,41\] receptors reduce oxidation parameters in cardiovascular system including heart. There are contradictory reports regarding modulation of oxidative stress by TRPV1 and GPR55 receptors \[30,31\].

Although CBD probably does not work via endocannabinoid receptors directly, it may act through augmentation of endocannabinoid tone \[42\]. CBD inhibits fatty acid amide hydrolase (FAAH) \[43\] and can interact with the anandamide membrane transporter \[44,45\] both of which may increase levels of endocannabinoids and related lipids. They may have positive effects and be used as a target in pharmacotherapy \[46\] but in some cases, can also exert untoward actions \[47\]. In this context, one should keep in mind that the FAAH inhibitor URB597 and hypertension may affect cardiac and plasma oxidative stress, endocannabinoid levels and lipid metabolism in a model-dependent manner \[48,49\].

The first aim of this study was to investigate whether chronic, unlike acute \[15\], administration of CBD reduces blood pressure (BP) and heart rate (HR) in rats with primary and secondary hypertension. Moreover, we studied whether CBD has an impact (ii) on the redox system, (iii) the endocannabinoid system in heart and plasma and (iv) free polyunsaturated fatty acids (PUFAs) and phospholipid PUFAs.

2. Results

2.1. General

As shown in Table 1 and Figure 1 (in which cardiovascular parameters were measured by the non-invasive method and telemetrically, respectively) SBP and DBP, registered before the first administration of CBD or its vehicle, were higher in SHR and DOCA rats than in the respective control animals (WKY and SHAM). HR tended to be lower in DOCA compared to SHAM rats and higher in SHR than in WKY when non-invasive registration was used (Table 1); in animals with telemetrical registration, HR was higher in WKY than in SHR (Figure 1). Two-week administration of CBD 10 mg/kg did not affect SBP, DBP and HR in normo- or hypertensive rats. The vehicle for CBD decreased
(or tended to decrease) HR by about 7%–8% in WKY and SHAM. Similarly, CBD reduced HR by about 10% in WKY (Table 1). We did not observe any cardiovascular effect of CBD during 24-h observation on the first and the last day of its administration (Figure 1).

**Figure 1.** Influence of cannabidiol (CBD) or its vehicle on systolic blood pressure (SBP), diastolic blood pressure (DBP) and heart rate (HR) in spontaneously hypertensive rats (SHR) and their normotensive controls (WKY). Parameters were measured telemetrically every hour (shown for DAY 1 and 14 in the intermediate and bottom panels, respectively) and means for each day were determined (shown in the top panels). CBD (10 mg/kg) was injected i.p. once daily for 14 days (control animals received the vehicle for CBD instead); first and final injection on DAY 1 and 14, respectively. Data are expressed as the median with interquartile range; \( n = 4 \). All values of SBP and DBP in SHR (in the presence of CBD or its vehicle) were higher than their respective values in WKY (\( p < 0.001 \)), and values of HR in SHR were lower than in WKY (\( p < 0.001 \) with 2 exceptions at the 14th day: WKY vs. SHR \( p < 0.05 \) and WKY + CBD vs. SHR + CBD: \( p < 0.01 \)).

The body weight of SHR and DOCA rats was lower than that of WKY and SHAM rats, respectively, both before the first and after the final administration of the vehicle of CBD (Table 1). When animals were considered which had received CBD instead, differences were smaller or did not occur at all (Table 1).
Table 1. Influence of cannabidiol (CBD) or its vehicle on systolic blood pressure, heart rate and body weight in conscious spontaneously (SHR) and deoxycorticosterone (DOCA) hypertensive rats and their respective normotensive controls (Wistar-Kyoto (WKY) and sham-operated (SHAM) rats).

| Group   | Treatment | n  | Systolic Blood Pressure (mmHg) | Heart Rate (Beats/min) | Body Weight (g) |
|---------|-----------|----|--------------------------------|------------------------|-----------------|
|         |           |    | Before the First Dose of CBD or Its Solvent | 24 h After the Final Dose of CBD or Its Solvent | Before the First Dose of CBD or Its Solvent | 24 h After the Final Dose of CBD or Its Solvent |
| WKY     | vehicle   | 7  | 111 [95;122]                    | 98 [91;105]           | 341 [321;366]   | 341 [304;326]   | 342 [329;361]   | 360 [367;400]   |
| WKY     | CBD       | 7  | 101 [87;124]                    | 92 [87;107]           | 334 [313;357]   | 300 [298;321]   | 325 [325;340]   | 360 [353;390]   |
| SHR     | vehicle   | 7  | 178 [161;199] ***               | 174 [164;187] ***     | 359 [351;369]   | 367 [365;387] ** | 290 [282;292] *** | 300 [296;326] ***|
| SHR     | CBD       | 7  | 184 [172;192] $$S$$            | 172 [170;176] $$S$$   | 355 [352;397]   | 358 [344;380]  $$S$$ | 315 [300;325]   | 328 [313;340]  $$S$$ |
| SHAM    | vehicle   | 7  | 122 [111;132]                   | 120 [118;131]          | 363 [351;371]   | 336 [331;340]   | 294 [270;304]   | 320 [302;345]   |
| SHAM    | CBD       | 7  | 111 [100;129]                   | 121 [109;134]          | 354 [333;374]   | 349 [328;371]   | 278 [256;294]   | 322 [286;340]   |
| DOCA    | vehicle   | 7  | 163 [111;175]                   | 175 [160;190] ***     | 314 [303;360]   | 324 [312;352]   | 275 [233;276]   | 280 [252;321]   |
| DOCA    | CBD       | 6  | 150 [143;174] $$S$$             | 173 [150;189] $$S$$   | 327 [315;354]   | 314 [301;348]   | 280 [271;298]   | 303 [297;312]   |

Cardiovascular parameters were measured by the non-invasive tail-cuff method. CBD (10 mg/kg) or its vehicle were injected intraperitoneally once daily for 14 days. Data are expressed as the medians with interquartile range; $^S$ $p < 0.05$; $$p < 0.01$; $$$p < 0.001$ significantly different from the respective values obtained in normotensive groups receiving vehicle for CBD (*) or CBD ($^S$) or recorded before the first dose of the compound ($^!$).
2.2. Influence of Hypertension and CBD on the Endocannabinoid System

As shown in Figure 2, 2-AG showed the highest level among the endocannabinoids; values in the heart of normotensive WKY and SHAM rats (expressed as median (interquartile range)) amounted to 5.0 (4.7;7.5) \( (n = 7) \) and 1.3 (1.1;1.7) nmol/g tissue \( (n = 7) \), respectively. The levels of the best-known endocannabinoid AEA were relatively low, i.e., about 220 and 75 times lower than the respective values of 2-AG in WKY and SHAM rats. Although levels of PEA, LEA and OEA were higher than those of AEA, they were still lower than 2-AG levels by factors of about 15, 40 and 60 (WKY) and 5, 20 and 25 (SHAM), respectively. HEA, DEA and DGLEA showed the lowest levels, i.e., about 1–3 pmol/g tissue in all cases.

![Figure 2](image-url)

**Figure 2.** Influence of hypertension and cannabidiol (CBD) or its vehicle on endocannabinoid levels in heart isolated from spontaneously (SHR; A) and deoxycorticosterone-salt (DOCA; B) hypertensive rats and their normotensive controls (WKY (A) and SHAM (B), respectively). CBD (10 mg/kg) was injected i.p. once daily for 14 days; the respective controls received the vehicle of CBD instead; hearts were prepared 24 h after the final injection. Data are expressed as the median with interquartile range; \( n = 5-7; * / # p < 0.05; ** / ## p < 0.01; *** p < 0.001 \) significantly different from WKY or SHAM (*) and from SHR or DOCA (#). AEA—anandamide; 2-AG—2-arachidonoylglycerol; PEA—palmitoyl ethanolamide; LEA—linolenoyl ethanolamide; OEA—oleoyl ethanolamide; SEA—stearoyl ethanolamide; DHEA—docosahexaenoyl ethanolamide; POEA—palmitoleoyl ethanolamide; HEA—homo-\( \gamma \)-linolenyl ethanolamide; DEA—docosatetraenoyl ethanolamide; DGLEA—dihomo-\( \gamma \)-linolenoyl ethanolamide; FAAH—fatty acid amide hydrolase; MAGL—monoacylglycerol lipase; m-NA—m-nitroaniline; DTNB—5,5′-dithiobis-2-dinitrobenzoic acid.

Hypertension modified cardiac endocannabinoid levels in a model-dependent manner (Figure 2). Thus, they mostly decreased (2-AG by 24% and LEA by 51%) or tended to decrease (AEA, DHEA...
POEA, HEA, DEA and DGLEA) in SHR in comparison to WKY. In contrast, they increased (2-AG by 158%, LEA by 97%, OEA by 180%, DHEA by 56%, DEA by 40% and DGLEA by 54%) in DOCA hearts in comparison to SHAM. Chronic CBD administration did not modify the cardiac endocannabinoid levels in SHR and WKY. However, it decreased levels of 2-AG (−57%), OEA (−28%), DEA (−15%) and DGLEA (−20%) in hearts of DOCA but increased PEA (+68%) and HEA (+80%) levels in SHAM rats.

In both models of hypertension, we observed increases in FAAH activities in comparison to the respective normotensive control (+77% in SHR and +115% in DOCA), but no changes in MAGL activities (Figure 2). CBD did not modify MAGL activity but decreased FAAH activity in DOCA and tended to do so in SHR. Moreover, it increased FAAH activity in WKY.

Plasma AEA levels were about 1 nM both in WKY and SHAM rats (Figure 3). For both control groups, levels were higher (PEA and OEA by about 10 and 5 times, respectively) and similar (SEA) when compared to the values of AEA. In contrast to SHR hearts, the plasma endocannabinoid levels increased (SEA +38%, DGLEA +63%) or tended to increase (OEA, POEA DHEA and DEA) in comparison to WKY. In DOCA, plasma levels increased (AEA +38% and LEA +113%), decreased (OEA −29% and DHEA −41%) or tended to decrease (POEA and DEA) in comparison to SHAM rats.

Chronic CBD administration decreased plasma levels of SEA (−25%), HEA (−23%) and DGLEA (−39%) in SHR and AEA (−29%) and LEA (−44%) in DOCA or tended to do so in the case of PEA, OEA and LEA in SHR (Figure 3). Moreover, CBD decreased plasma levels of HEA (−20%) in WKY and
increased the plasma level of AEA (+33%) in SHAM or tended to decrease levels of AEA, OEA, POEA, DEA and DGLEA in WKY and levels of POEA and DHEA in SHAM.

In the cardiac left ventricle, cannabinoid CB₁ and CB₂ receptor density decreased but GPR18 and GPR55 density increased in SHR compared to WKY. In DOCA, compared to SHAM, only a decrease in CB₁ density and an increase in GPR18 density occurred. The increase in GPR18 receptor density was particularly marked in both hypertension models (by about 60%) (Figure 4). Chronic CBD administration decreased GPR55 density in SHR and tended to decrease CB₁ and GPR18 receptor density in SHR and CB₁, CB₂ and GPR18 receptor density in DOCA. Moreover, CBD decreased CB₁ and tended to reduce GPR18 receptor density in WKY. TRPV1 receptor density was not modified by hypertension or CBD treatment (Figure 4).

**Figure 4.** Influence of hypertension and cannabidiol (CBD) or its vehicle on expression of receptors in left ventricle isolated from spontaneously (SHR; A) and deoxycorticosterone-salt (DOCA; B) hypertensive rats and their normotensive controls (WKY (A) and SHAM (B), respectively). CBD (10 mg/kg) was injected i.p. once daily for 14 days; the respective controls received the vehicle of CBD instead. Hearts were prepared 24 h after the final injection. Receptor protein was determined by Western blots and given as fraction of the value in the normotensive control (first of the four columns). Images obtained using stain-free gel technology that allows for total protein visualization and quantification are shown as a loading control (Loading). Data are expressed as the median with interquartile range; n = 5–6; * p < 0.05; ** p < 0.01; ***/### p < 0.001 significantly different from WKY or SHAM (*) and from SHR or DOCA (#).
2.3. Influence of Hypertension and CBD on Oxidative Stress

As shown in Figure 5, activities of cardiac antioxidant principles underwent slight changes in hypertension. Thus, in comparison to the respective normotensive tissues, GPx activity increased or tended to increase in DOCA (+48%) and SHR, respectively, whereas GSR activity decreased (−12%) in DOCA heart. However, SOD and CAT were not altered in SHR and DOCA. The cardiac vitamin A (but not E) level increased in SHR (+198%), whereas the levels of both antioxidant vitamins did not change in the heart of DOCA rats. Levels of glutathione and glutathione disulfide were altered by both models of hypertension in the same way: GSH decreased in the heart of SHR (−30%) and DOCA (−42%) and GSSG increased (+82% and +149%, respectively). Products of lipid peroxidation increased in the heart of SHR (4-HNE +45% and 4-HHE +86%) and DOCA rats (MDA +82%). Changes in cardiac CO groups were neither observed in SHR nor in DOCA.

Figure 5. Influence of hypertension and cannabidiol (CBD) or its vehicle on activity/level of oxidative stress parameters in heart isolated from spontaneously (SHR; A) and deoxycorticosterone-salt (DOCA; B) hypertensive rats and their normotensive controls (WKY (A) and SHAM (B), respectively). CBD (10 mg/kg) was injected i.p. once daily for 14 days; the respective controls received the vehicle of CBD instead. Hearts were prepared 24 h after the final injection. Data are expressed as the median with interquartile range; n = 5–7; * p < 0.05; **/### p < 0.01; ***/##### p < 0.001 significantly different from WKY or SHAM (*) and from SHR or DOCA (#). CAT—catalase; GSR—glutathione-disulfide reductase; GPx—glutathione peroxidase; SOD—superoxide dismutase; GSH—glutathione; GSSG—glutathione disulfide; CO groups—protein carbonyl groups; MDA—malondialdehyde; 4-HNE—4-hydroxynonenal; 4-HHE—4-hydroxyhexenal.

Chronic CBD administration was associated mainly with a decrease in cardiac levels of the products of lipid peroxidation (Figure 5). The effect was significant for 4-HHE (−37%) in SHR and for MDA (−63%) in DOCA and a tendency was observed in the case of 4-HNE in SHR and DOCA. Moreover, cardiac levels of vitamin A (−77%) and E (−64%) were reduced in SHR or tended to be so in DOCA. In addition, the level of GSH was increased (+29%) and that of GSSG was decreased (−29%) in DOCA; in SHR, both parameters tended to be altered. Other cardiac parameters of oxidative stress were not modified by CBD in hypertensive rats (Figure 5). In normotensive rats, CBD increased levels of...
vitamin A in SHAM (+41%) and unexpectedly levels of products of lipid peroxidation (MDA (+42%), 4-HNE (+61%) and 4-HHE (+86%)) in WKY but not in SHAM (Figure 5).

Similarly to the heart, plasma activities of antioxidant principles underwent only small changes in hypertension in comparison to normotension (Figure 6). Thus, only in SHR SOD (+3%) and GPx activities increased or tended to increase, respectively. The level of vitamin E decreased in SHR (−65%), but was not modified in DOCA. Levels of vitamin A did not undergo changes in either hypertension model. Levels of GSH and GSSG were modified in a comparable way in SHR and DOCA, i.e., they decreased and increased by about 45% and 175%, respectively. Levels of MDA (but not of other products of lipid peroxidation) increased by about 65% both in SHR and DOCA. The CO group level increased in SHR (+46%) and tended to do so in DOCA.

**Figure 6.** Influence of hypertension and cannabidiol (CBD) or its vehicle on activity/level of oxidative stress parameters in plasma isolated from spontaneously (SHR; A) and deoxycorticosterone-salt (DOCA; B) hypertensive rats and their normotensive controls (WKY (A) and SHAM (B), respectively). CBD (10 mg/kg) was injected i.p. once daily for 14 days; the respective controls received the vehicle of CBD instead. Plasma was obtained 24 h after the final injection. Data are expressed as the median with interquartile range; n = 5–7; */# p < 0.05; **/## p < 0.01; ***/### p < 0.001 significantly different from WKY or SHAM (*) and from SHR or DOCA (#). GSR—glutathione-disulfide reductase; GPx—glutathione peroxidase; SOD—superoxide dismutase; GSH—glutathione; GSSG—glutathione disulfide; CO groups—protein carbonyl groups; MDA—malondialdehyde; 4-HNE—4-hydroxynonenal; 4-HHE—4-hydroxyhexenal.

Chronic CBD administration increased plasma levels of vitamin E (+39%) in DOCA. GSH was increased (+34%) in DOCA and tended to be increased in SHR whereas GSSG tended to decrease in either hypertensive rat model. For antioxidant enzyme activities only tendencies towards a decrease of GPx (SHR and DOCA) and GSR (DOCA) were observed. However, CBD reduced plasma levels of CO groups (-14%) in SHR and of MDA (-45%) in DOCA. Moreover, it tended to decrease plasma levels of 4-HNE (-32%) in SHR and 4-HHE (-37%) in DOCA. Unexpectedly it also increased plasma levels of MDA (+38), 4-HNE (+101%) and 4-HHE (+62) in WKY and of 4-HHE (+39%) in SHAM (Figure 6).
2.4. Influence of Hypertension and CBD on Lipids

The effect of hypertension on free fatty acids and phospholipids was studied in cardiac tissue (Figure 7A,B) and plasma (Figure 7C,D). Among FFA, hypertension increased and decreased levels of AA (+50%) and of LA (−51%) in the heart of DOCA, respectively, but did not modify cardiac levels of FFA in SHR and plasma levels both in SHR and DOCA. In the fraction of phospholipids, hypertension increased plasma levels of AA (+25%) and DHA (+48%) and tended to increase plasma levels of LA in DOCA. On the other hand, it decreased cardiac levels of DHA (−25%) in SHR and LA (−28%) in DOCA and plasma levels of LA (−16%), AA (−42%) and DHA (−51%) in SHR.

Chronic CBD administration increased cardiac levels of FFA LA (+67%) and FFA AA (+32%) in SHR and FFA LA (+100%) in DOCA but decreased FFA AA in the heart of DOCA (−39%) and in the plasma of SHR (−32%). In normotensive controls, CBD increased or tended to increase cardiac levels of FFA LA (+60%) and FFA AA (+28%) in WKY and cardiac levels of FFA LA (+123%) and FFA AA (+82%) in SHAM. On the other hand, CBD decreased plasma levels of FFA AA (−26%) and PH DHA (−15%) in WKY and FFA LA (−36%) and FFA AA (−68%) in SHAM.

Figure 7. Influence of hypertension and cannabidiol (CBD) or its vehicle on levels of phospholipids (PH) and free fatty acids (FFA) in heart (A,B) and plasma (C,D) isolated from spontaneously (SHR; A,C) and deoxycorticosterone-salt (DOCA; B,D) hypertensive rats and their normotensive controls (WKY (A,C) and SHAM (B,D), respectively). CBD (10 mg/kg) was injected i.p. once daily for 14 days; the respective controls received the vehicle of CBD instead. Hearts were prepared and plasma was obtained 24 h after the final injection. Data are expressed as the median with interquartile range; n = 5–7; * p < 0.05; **/## p < 0.01; *** p < 0.001 significantly different from WKY or SHAM (*) and from SHR or DOCA (#). LA—linoleic acid; AA—arachidonic acid; DHA—docosahexaenoic acid.
3. Discussion

3.1. General

This study shows that chronic CBD administration did not modify BP and HR in rats with primary (SHR) and secondary (DOCA-salt) hypertension in spite of the reduction of cardiac and plasma oxidative stress. It inhibited the FAAH activity in both hypertension models, but had opposite effects on cardiac levels of various endocannabinoids and endocannabinoid-related lipids (decrease in SHR vs. increase in DOCA). Unexpectedly, CBD increased lipid peroxidation in normotensive controls and this alteration may lead to untoward effects.

Our study is based on rats with primary (SHR; the most frequently studied genetic hypertensive model [17] and secondary hypertension (DOCA-salt [17]; because a salt-rich diet is one of the main lifestyle factors leading to hypertension). WKY and SHAM served as the respective normotensive controls and allowed us to detect potential undesirable effects of CBD. We used CBD at 10 mg/kg for 14 days since its chronic i.p administration had a beneficial effect on cardiovascular tissues (including decreases in oxidative stress) from diabetic (7 days; [50]) and septic rats (9 days; [28]) and from mice with diabetic (11 weeks; [22]) and doxorubicin-induced cardiomyopathy (5 days; [26]) and experimental autoimmune myocarditis (46 days; [51]). Note that CBD was used at ~10 mg/kg also in several acute experiments [10,13,15].

To examine the mechanism(s) of CBD action in hypertension, we considered parameters of oxidative stress (e.g., GSH and GSSG), of the endocannabinoid system (ECS) and of other components of lipid metabolism. With respect to the ECS, the two endocannabinoids AEA and 2-AG and the endocannabinoid-related lipids POEA, HEA, DEA, DGLEA, PEA, LEA, OEA, SEA and DHEA were determined [52]. The latter five and AEA are degraded by FAAH [53,54]. FAAH and MAGL activities and CB1, CB2, GPR18, GPR55 and TRPV1 receptor expression have been quantified as well. With respect to lipid metabolism free polyunsaturated fatty acids (PUFAs) including AA, LA and DHA and phospholipid PUFAs were determined.

The present results confirm and extend our previous observations [49] that in the heart, 2-AG has the highest concentration among the 11 endocannabinoids or endocannabinoid-related lipids. Its levels in WKY and SHAM were 220 and 75 times higher than the concentration of the best-known endocannabinoid AEA, respectively. This is an important observation in the light of recent publications that 2-AG in vivo worsens heart function after acute myocardial infarction [47], increases the severity of the cerebral blood flow deficit [55] or promotes atherogenesis [56] on the one hand but ameliorates inflammatory stress-induced insulin resistance in cardiomyocytes on the other [57]. Unfortunately, there are only limited data regarding the cardiovascular effects of endocannabinoids other than 2-AG and AEA. Thus, chronic administration of PEA (30 mg/kg for 5 weeks) or OEA (5 mg/kg for 7 days) improved rat myocardial function in doxorubicin-induced heart failure [58] and decreased BP and protected against kidney injury in SHR via inhibition of oxidative stress [59].

3.2. Effect of Hypertension

SBP was elevated in primary and secondary hypertension of animals in which BP was determined by the non-invasive (both models) and/or telemetric (SHR) method. With respect to SHR, both methods revealed an identical SBP (of about 170 mmHg) whereas HR differed. Compared to WKY, HR in SHR was higher or lower when the tail-cuff method or telemetry was used, respectively. A significant influence of restraint on HR determination by the non-invasive method (higher HR in SHR than in WKY) was described earlier [60]. In our previous study [49], DOCA-salt treatment over a period of 6 weeks resulted in a much higher SBP in DOCA than in SHR (about 220 vs. 185 mmHg, respectively). To avoid such a difference in the current study we reduced DOCA-salt treatment to 4 weeks (prior to the onset of the CBD treatment) which resulted in comparable SBP values on the final day of experiments in SHR and DOCA treated with vehicle (about 170 mmHg).
The elevated BP was connected with a redox imbalance, similarly to our previous papers [48,49]. Thus, GSSG increased and GSH decreased in heart and plasma of both hypertensive models. Other markers, including increased MDA, 4-HNE, 4-HHE and oxidative protein modifications (CO) occurred only in one hypertensive model and/or only in heart or plasma. In SHR (but not in DOCA) a decrease in plasma vitamin E and, contrarily to expectation, an increase in cardiac vitamin A was observed.

Cardiac FAAH activities increased in both models of hypertension like in our previous study [49]. Moreover, the present results show that in addition to AEA and 2-AG [49] also LEA, DHEA, POEA, HEA, DEA and DGLEA decreased (or tended to decrease) in the heart of SHR whereas in the heart of DOCA not only 2-AG but also LEA, OEA, DHEA, DEA and DGLEA increased. On the rule, changes in plasma endocannabinoid levels were similar (but less pronounced) with the exception that in DOCA OEA and DHEA levels decreased. Since numerous endocannabinoid-like compounds were changed in hypertensive animals, future studies should clarify whether these compounds affect the cardiovascular system. As suggested previously [49], the enhanced levels of AEA, LEA, OEA and SEA (all of which are degraded by FAAH) in spite of higher FAAH activity might indicate that in the heart of the DOCA group endocannabinoid synthesis is favoured [61] and/or AEA transporter activity is decreased [62].

Cardiac CB1 receptor density decreased and GPR18 receptor density increased in both hypertension models whereas CB2 receptor density decreased in SHR only. The results obtained with cannabinoid and cannabinoid-like receptors only partially conform to those in our previous study [49]; differences may result from the fact that (i) receptor densities were determined in left ventricle vs. whole heart, respectively, and (ii) the time period for induction of DOCA-salt hypertension lasted for 4 vs. 6 weeks, respectively. As mentioned in the Introduction, stimulation of CB1 receptors enhances oxidative stress, whereas CB2 and GPR18 receptors have an opposite effect. Accordingly, the hypertension-induced enhancement in oxidative stress does not fit well to the changes in receptor densities since in both hypertension models, a decreased pro-oxidative receptor (CB1) was connected with an increased anti-oxidative receptor (GPR18).

The hypertension-related changes in lipids other than the endocannabinoids were dependent on the hypertension model. Similarly to endocannabinoids, mainly decreases in phospholipids (cardiac PH DHA and plasma PH LA, AA and DHA) were obtained in SHR. By contrast, in DOCA plasma phospholipid AA and DHA and cardiac FFA AA increased whereas cardiac FFA LA and PH LA decreased. Differences in particular cardiac and plasma PUFAs between SHR and DOCA were also observed in our previous study [49].

3.3. Effect of Chronic CBD in Hypertensive Animals

Chronic administration of CBD failed to modify BP and HR in both models of hypertension; this is in harmony with our previous study [15], in which a single CBD administration had no effect. One may argue that the dose of CBD was too low or the duration of its application too short; however, both parameters conformed to the conditions chosen in previous studies in which CBD had a beneficial effect (see 3.1). One may also argue that the effectiveness of CBD was lost during the study; however, FAAH (which is inhibited by CBD; [43]) was still reduced (DOCA) or tended to be reduced (SHR) 24 h after the final dose of CBD.

How can we explain the lack of an antihypertensive effect of CBD in hypertension? CBD does not only possess vasodilatory activity mainly shown on isolated vessels (for review, see [9]) but, according to our recent study in pithed SHR and WKY [15], at an intravenous dose of 10 mg/kg, it also exhibits sympathomimetic effects. Opposite cardiovascular effects might be at least partially responsible for the lack of a hypotensive effect of CBD in our study.

Hypertension is associated with an enhancement of oxidative stress ([16]; current study) and CBD is known for its antioxidant properties (for literature, see Introduction). Indeed, chronic CBD administration led to increases in GSH and decreases in GSSG both in heart and plasma in both hypertension models. Moreover, CBD counteracted parameters of lipid peroxidation including the enhanced cardiac and/or plasma 4-HNE and 4-HHE in SHR and the enhanced cardiac and plasma
MDA and 4-HNE in DOCA. Moreover, CBD reduced the enhanced concentration of carbonyl groups in plasma. The increase in plasma vitamin E in DOCA was another beneficial effect of CBD although, contrarily to expectation, cardiac levels of vitamin A and E in SHR decreased.

The endocannabinoid system is overactivated in hypertension [63]. Chronic CBD administration failed to modify the densities of cannabinoid and cannabinoid-like receptors (the only exception was the decrease in cardiac GPR55 levels in SHR) but decreased cardiac levels of 2-AG, OEA, DEA and DGLEA in DOCA and plasma levels of AEA and LEA in DOCA and of SEA, HEA and DGLEA in SHR. On the rule, CBD reduced the levels of those ECBs the concentrations of which were enhanced in hypertension. Accordingly, CBD failed to modify cardiac endocannabinoid levels in SHR since only decreases were obtained. Surprisingly, the decreases in ECBs were observed in spite of the reduced FAAH activity. Due to the vasodilatory and/or hypotensive effects of AEA and OEA [64,65], the CBD-induced decrease in AEA and OEA levels appears to be unfavourable in hypertension whereas the decrease in cardiac 2-AG concentration might be beneficial because worsening of cardiac function by 2-AG has been described recently [47].

The effect of CBD on PUFAs was again dependent on the hypertension model and on the level of PUFAs in hypertension. Hypertension-induced increases in PUFAs were observed in DOCA only and CBD decreased the cardiac FFA AA and tended to decrease the plasma PH AA and DHA.

### 3.4. Effect of Chronic CBD in Normotensive Animals

CBD is generally recognized as a safe drug [66]. Surprisingly, we have obtained some unexpected effects of chronic CBD administration in normotensive rats. The most untoward influence was the enhancement of lipid peroxidation documented by increases of heart and plasma MDA, 4-HNE and 4-HHE in SHAM and, in addition, of plasma 4-HHE in WKY. These alterations were connected with a decrease in cardiac (antioxidant) GPR18 receptor density in WKY and an increase (or the tendency of an increase) in cardiac FFA LA and FFA AA in WKY and SHAM. Our findings, although not unequivocal (since in plasma decreases in FFA AA (WKY and SHAM), FFA LA (SHAM) and HEA (WKY) occurred), are reminiscent of a previous study [28], in which CBD (10 mg/kg) administered to rats for 9 days led to an increase in carbonyl groups in lung and liver.

### 3.5. Limitations of the Study

The present investigation, in which a blood pressure-lowering effect of CBD could not be shown despite its antioxidant effect, was restricted to two models of hypertension studied in male rats. The possibility has to be considered that an antihypertensive effect of CBD was missing for the following reasons. (1) The duration of CBD administration might have been too short or its dose too low and an increase in the duration of its application or the use of a higher dose might have led to more evident changes, especially of the markers of oxidative stress and probably also of the level of blood pressure. (2) In our previous paper, the chronic administration of the FAAH inhibitor URB597 decreased BP in DOCA but not in SHR, in which BP before the first dose of URB597 was about 220 vs. 185 mmHg, respectively [67]. As mentioned above, in the present study, the period for the induction of DOCA hypertension was reduced from 6 to 4 weeks, which resulted in lower SBP (about 170 mmHg). We cannot exclude that the lack of effect of CBD is related to the shorter induction time and the lower blood pressure level in DOCA. (3) We used DOCA as a secondary model of hypertension. The possibility has to be considered that a model of hypertension connected with changes in the renin-angiotensin-aldosterone system (RAAS) (the activity of which is modified by the endocannabinoid system) might have revealed a blood pressure-lowering effect of CBD. Interestingly, opposite effects of a CB1 receptor antagonist have been found in the cardiovascular system of SHR and of rats with a RAAS-dependent hypertension (for review, see [63]). (4) There are gender-specific differences in hypertension [17,68] and it would be interesting to extend our experiments to female rats. (5) There is no ideal animal model of human hypertension. However, the advantage of rat genetic models of hypertension is their similarity to the BP/hypertension phenotypes observed in patients.
and that SHR responds to the antihypertensive effects of almost all classes of drugs approved for treatment of hypertension. The DOCA-salt model connected with unilateral nephrectomy provides a reliable animal model that can develop severe hypertension with some features of human low-renin hypertension [17].

The mechanism of action for the antioxidant effect of CBD was not determined. This is not a trivial task since the compound has many different molecular effects [1,3,4,29]. The fact that CBD influenced components of the ECS in the two hypertension models in an opposite manner suggests that a direct antioxidant effect is more likely than an indirect ECS-based mechanism. Finally, although the antioxidant effect of CBD was not associated with an antihypertensive effect it would be interesting to examine the influence of CBD on other cardiovascular parameters, e.g., arrhythmia risk, in future studies. Thus, oxidative stress was shown to be involved in cardiac electrical and structural remodelling [69] and in the pathophysiology of atrial fibrillation [70]. The use of ECG predictors of arrhythmia risk would be a benefit in such studies [69,71].

4. Materials and Methods

4.1. Animals

All procedures and experimental protocols were performed in accordance with the European Directive (2010/63/EU) and with the approval of the local Animal Ethics Committee in Olsztyn (Poland) (Approval code: 80/2017; approval date: 28 November 2017). Rats were obtained from the Center for Experimental Medicine of the Medical University of Białystok (Poland). They had free access to chow and water and were kept under a 12:12 h light-dark cycle. Experiments were performed on male rats with spontaneous (SHR) and secondary DOCA-salt hypertension.

4.2. Experimental Groups and Protocol

DOCA-salt hypertension was induced in Wistar rats. After unilateral nephrectomy of 5-6 week-old animals and one week of recovery, deoxycorticosterone acetate (DOCA) was injected s.c. (25 mg/kg in 0.4 mL DMF/kg) twice a week for 28 days. During DOCA administration, drinking water was replaced with 1% saline water. Control group for DOCA were sham-operated (unilaterally nephrectomised) (SHAM) Wistar rats. They received s.c. DMF (DOCA vehicle) twice weekly for 4 weeks and tap water for drinking.

Animals were randomly divided into experimental groups: (1) hypertensive DOCA rats, (2) respective normotensive control SHAM rats, (3) hypertensive SHR rats and (4) respective normotensive WKY rats. All animal groups were age-matched at the beginning of CBD treatment (8–9 weeks old). The body weight of the rats is shown in Table 1.

4.3. Chronic CBD Administration

One part of every hypertensive and normotensive group were injected i.p. with CBD (10 mg/kg) every 24 h for 14 days. The other part received CBD vehicle (ethanol, Tween 80, 0.9% NaCl—3:1:16; 1 mL/kg).

4.4. Determination of Cardiovascular Parameters in Conscious Rats

Systolic blood pressure (SBP) and heart rate (HR) were measured using non-invasive tail-cuff method with Non-Invasive Blood Pressure Controller (ADInstruments, Sydney, Australia) before first dose of CBD or its vehicle and after 7 and 14 days of experiment (24 h after last dose of compounds).

SBP, diastolic blood pressure (DBP) and HR were also measured telemetrically in SHR and WKY rats, as described previously [15]. Briefly, after pentobarbitone sodium (300 µmol/kg; i.e., ~70 mg/kg; i.p.) anesthesia, telemetry transmitters (HD-S10, Data Sciences International, Saint Paul, MN, USA) were implanted into the femoral artery. Rats were allowed to recover for 1 week before measurements.
4.5. Tissue Preparation for Biochemical Examinations

Twenty-four hours after the last dose of CBD or its vehicle rats were anesthetized with pentobarbitone sodium (300 µmol/kg; i.p.) to collect blood and heart. Blood samples were obtained by left ventricle puncture and collected into EDTA tubes. Plasma separation from whole blood was carried out by centrifugation at 2000× g for 5 min.

Hearts were perfused with 0.9% saline and cut lengthwise into two halves with equal size and quality. The first part of the tissue was snap-frozen with liquid nitrogen and stored at −80 °C. The second part was homogenized. 10% homogenates in 0.9% saline were centrifuged at 20000× g for 15 min at 4 °C.

4.6. Biochemical Studies

4.6.1. Determination of Endocannabinoids

Anandamide (AEA), 2-arachidonoylglycerol (2-AG), palmitoyl ethanolamide (PEA), linolenoyl ethanolamide (LEA), oleoyl ethanolamide (OEA), stearoyl ethanolamide (SEA), docosahexaenoyl ethanolamide (DHEA), palmitoleoyl ethanolamide (POEA), homo-γ-linolenyl ethanolamide (HEA), docosatetraenoyl ethanolamide (DEA) and dihomo-γ-linolenoyl ethanolamide (DGLEA) were determined using modified ultrahigh performance liquid chromatography-tandem mass spectrometry (UPL-CMS/MS) by the Lam method [72]. Octadeuterated endocannabinoids: AEA-d₈, 2-AG-d₈ and OEA-d₄ [73] as internal standards were added into the tissue lysates and all cannabinoids were isolated using a solid phase extraction (OASIS HLB 3cc). UPLC–MS/MS analysis was carried out using an Nexera X2 Shimadzu UPLC system with a Zorbax Extend C18 column (2.1 mm×150 mm, 1.8 mm, Agilent, Santa Clara, CA, USA) and interfaced with a Shimadzu 8060 triple quadrupole mass spectrometer with electrospray ionization source (ESI). The samples were analyzed in positive-ion mode using multiple reaction monitoring (MRM). Transitions of the precursor to the product ion was as follows: m/z 348.3 → 62.15 for AEA, m/z 379.3 → 287.25 for 2-AG, m/z 300.3 → 62.00 for PEA, m/z 324.0 → 62.00 for LEA, m/z 326.0 → 62.00 for OEA, m/z 328.0 → 62.00 for SEA, m/z 372.0 → 62.00 for DHEA, m/z 298.0 → 62.00 for POEA, m/z 314.5 → 62.00 for HEA, m/z 376.0 → 62.00 for DEA and m/z 350.0 → 62.00 for DGLEA.

4.6.2. Determination of FAAH and MAGL Activity

Fatty acid amide hydrolase (FAAH) (EC.3.5.1.99) activity was measured in the homogenate of heart tissue prepared in 20 mM Tris, containing 10% glycerol, 150 mM NaCl, and 1% Triton X-100, pH 7.8 at 4 °C. Following centrifugation (1000× g), 20 µL of the supernatant was added to 175 µL of reaction buffer (125 mM Tris, pH 9.0, and 1 mM EDTA) and 17 µM of FAAH substrate, decanoyl m-nitroaniline. Formation of m-nitroaniline (m-NA) was determined at 410 nM [74]. Specific enzyme activity was expressed in nmoles of m-NA/min/mg protein.

Monoacylglycerol lipase (MAGL) (EC.3.1.1.23) activity was measured in the homogenate of heart tissue prepared in 20 mM Tris, 320 mM sucrose and 1mM EDTA, pH 8.0. Heart supernatant was obtained from the soluble fraction after spinning the homogenate at 1000× g for 15 min. A reaction mixture containing heart supernatants, 10 mM Tris, 1 mM EDTA pH 7.2 was pre-incubated at 4 °C for 15 min. After addition of arachidonoyl-1-thio-glycerol (A-1-TG), the mixture was incubated at 37 °C for 5 min and after refrigerating to room temperature, 1 mM DTNB was added. After 3 min, the formation of TNB was determined at 412 nm. Specific enzyme activity was expressed in nmoles of TNB/min/mg of protein [75].

4.6.3. Western Blot Analysis

A routine Western blotting procedure was used to examine protein expression, as has been described previously [67], except using stain-free technology [76]. Briefly, samples from the left ventricles were homogenized in radioimmunoprecipitation assay (RIPA) buffer containing a cocktail
of protease and phosphatase inhibitors. In addition, protein concentration was measured using the bicinchoninic acid method (BCA) with bovine serum albumin (BSA) as a standard. Subsequently, homogenates were reconstituted in Laemmli buffer. The same amounts of protein (30 µg) were loaded on Criterion™ TGX Stain-Free Precast Gels (Bio-Rad, Hercules, CA, USA). After electrophoresis, proteins were transferred onto PVDF (polyvinylidene difluoride) membranes using Trans-Blot Turbo Transfer System (Bio-Rad, Hercules, CA, USA) and stain-free blot image was taken (ChemiDoc XRS; Bio-Rad, Hercules, CA, USA) for total protein measurement in each sample lane. Next, the membranes were incubated overnight at 4 °C with the corresponding primary antibodies in appropriate dilutions: CB1 (1:500), CB2 (1:500), GPR18 (1:5000), GPR55 (1:1000) and TRPV1 (1:500). Thereafter, PVDF membranes were incubated with the appropriate secondary antibody conjugated to horseradish peroxidase (Cell Signaling Technology, Danvers, MA, USA). After adding a suitable substrate for horseradish peroxidase protein bands were detected using a ChemiDoc visualization system XRS (Bio-Rad, Hercules, CA, USA). The expression of selected target proteins was quantified using stain-free gels and total protein normalization method (Bio-Rad, Hercules, CA, USA).

4.6.4. Determination of Antioxidant Enzyme Activity

Catalase (CAT—EC.1.11.1.9) activity was measured in the homogenate of heart tissue by a spectrophotometric analysis (at 240 nm) of the rate of hydrogen peroxide decomposition, using a method published previously [77]. One unit of CAT is defined as the amount of the enzyme necessary to catalyze the decomposition of 1 µmol of hydrogen peroxide to water and oxygen within 1 min.

Glutathione reductase (GSR—EC.1.6.4.2) activity was measured according to the method of Mize and Langdon [78] by monitoring the oxidation of NADPH at 340 nm at a pH 7.4. One unit of GSR oxidized 1 µmol of NADPH/min at 25 °C and pH 7.4. Specific enzyme activity was expressed in units per mg of protein.

Glutathione peroxidase (GPx—EC.1.11.1.9) activity was assessed spectrophotometrically using the method of Paglia and Valentine [79]. GPx activity was assayed by measuring the conversion of NADPH to NADP+. One unit of GPx activity was defined as the amount of enzyme catalyzing the oxidation of 1 µmol NADPH/min at 25 °C and pH 7.4. Specific enzyme activity was expressed in units per mg of protein.

Superoxide dismutase (SOD—EC.1.15.1.1) activity was measured according the method by Sykes et al. [80]. The oxidation of epinephrine was performed in terms of the production of adrenochrome, which has a maximal absorption at 480 nm. One unit of SOD is defined as the amount of the enzyme that inhibits the rate of autoxidation of epinephrine by 50%.

4.6.5. Determination of Non-Enzymatic Antioxidant Level

Vitamin E and A were detected in the samples using high-performance liquid chromatography (HPLC) [81]. Extraction of vitamins was carried out using hexane. After removal, drying and dilution with ethanol, the hexane phase (50 µL) was injected on the column. UV detection at 294 nm for vitamin E and 298 nm for vitamin A were applied. The flow rate was 1 mL/min of methanol and water (95:5).

Glutathione (GSH) and glutathione disulfide (GSSG) were quantified using the capillary electrophoresis (CE) method of Maeso et al. [82]. Samples were sonificated in the Eppendorf tubes with 2 mL of a mixture containing ACN/H₂O (62.5:37.5, v/v) and centrifuged at 29,620×g for 10 min. The supernatant was immediately measured by CE. The separation was performed on a capillary with 47 cm total length (40 cm effective length) and 50 µm ID and was operated at 27 kV with UV detection at 200 ± 10 nm.
4.6.6. Determination of Protein Modifications

Protein oxidative modifications (carbonyl groups; CO groups) were determined according to the method published previously [83]. Carbonyl content was computed from peak absorption (370 nm) using 2,4-dinitrophenylhydrazine as a reagent.

4.6.7. Determination of Lipid Modifications

Malondialdehyde (MDA), 4-hydroxynonenal (4-HNE) and 4-hydroxyhexenal (4-HHE) were measured by GC/MSMS, as the O-PFBoxime-TMS derivatives, using modified method of Luo et al. [84]. Benzaldehyde-D6 as an internal standard was added to the tissue lysates and aldehydes were derivatized by the addition of O-(2,3,4,5,6-pentafluorobenzyl) hydroxyamine hydrochloride (0.05 M in PIPES buffer, 200 μL; incubation for 60 min at room temperature). Subsequently, samples were deproteinized by the addition of 1 mL of methanol and OPFB-oxime aldehyde derivatives were extracted by the addition of 2 mL of hexane. The top hexane layer was transferred into borosilicate tubes and evaporated under a stream of argon gas followed by the addition of N,O-bis(trimethylsilyl)trifluoroacetamide in 1% trimethylchlorosilane. A 1 μL aliquot was injected on the column. Derivatized aldehydes were detected in the selected ion monitoring (SIM) mode. The ions used for MDA/4-HNE/4HHE-PFB-TMS identification were m/z 204.0 and 178.0 for MDA; m/z 333.0 and 181.0 for 4-HNE; and 352.0 and 226.0 for 4-HHE respectively and m/z 307.0 for IS (benzaldehyde-D6) derivatives.

4.6.8. Determination of Fatty Acids

The concentration of the fatty acids AA, DHA and LA was determined by gas chromatography [85]. Lipid components were isolated from tissue lysates by extraction with chloroform/methanol mixture (2:1, v/v). Using TLC, total phospholipids were separated with the mobile phase heptane—diisopropyl ether—acetic acid (60:40:3, v/v/v). All lipid fractions were transesterified to fatty acid methyl esters (FAMEs) with boron trifluoride in methanol reagent under nitrogen atmosphere without previous separation from the layer. The FAMEs were quantified by gas chromatography with a flame ionization detector. Separation of FAME was carried out on a capillary column coated with Varian CP-Sil88 stationary phase and analyzed by gas chromatography with a flame ionization detector (FID) on a Clarus 500 Gas Chromatograph (Perkin Elmer, Waltham, MA, USA).

4.7. Statistical Analysis

The results are expressed as median values and interquartile range. Cardiovascular parameters were obtained from WKY and SHR rats in which blood pressure was recorded telemetrically (n = 4). Rats, in which blood pressure was determined by the tail-cuff method, served both for the registration of cardiovascular and biochemical parameters (WKY, SHR, SHAM, DOCA-salt). At the beginning, each group consisted of 7 rats. However, the final n was 5-7 because of (1) the death of one rat (DOCA + CBD) and/or (2) the exclusion of outliers (values deviating from the mean by more than plus/minus three standard deviations). All data were subjected to the Kolmogorov–Smirnov test to assess the distribution of values. If the data were normally distributed, parametric tests were done (paired Student’s t-test for comparison within group and one-way ANOVA with Bonferroni’s multiple comparison test for multiple groups). Data subjected to ANOVA were followed by Bonferroni’s post hoc tests only when the F value attained p < 0.05 and there was no significant inhomogeneity of variances. If the data were not normally distributed, a non-parametric test was performed (Wilcoxon test for comparison within group and Kruskal–Wallis test with Dunn’s post hoc test to compare multiple groups). Dunn’s post hoc test was only used when the Kruskal–Wallis test yielded a significant result (p < 0.05). A statistical analysis was performed using Graph Pad Prism 5 (GraphPad Software, La Jolla, CA, USA).
4.8. Drugs

(-)-cannabidiol (CBD) (THC-1073G-1) from THC Pharm, Frankfurt, Germany; ethanol (BA6420113) and natrium chloride (NaCl) (BA4121116) from POCH, Gliwice, Poland; Tween 80 (P1754), 11-deoxy cortisol acetate (DOCA) (D7000), N,N-dimethylformamide (DMF) (9227056), chloro-2,4-dinitro benzene (CDNB) (237329), butylated hydroxyloluene (BHT) (W218405) and 5,5′-dithiobis(2-nitrobenzoic acid) (DTNB) (D8130) from Sigma-Aldrich, Munich, Germany; pentobarbital sodium (5909991290153) from Biowet, Puławy, Poland; anandamide-d8 (AEA-d8) (390050), 2-arachidonoylglycerol-d8 (2-AG-d8) (362160), oleoyl ethanolamide-d4 (OEA-d4) (900552), decanoyl m-nitroanilinie (m-NA) (90349), arachidonoyl-1-thio-glycerol (A-1-TG) (10007904) from Cayman Chemical Company, Ann Arbor, MI, USA; CB1 antibody (ab23703), CB2 antibody (ab3561), GPR18 antibody (ab174835), GPR55 antibody (ab203663) from Abcam, Cambridge, UK; TRPV1 antibody (bs-1931R) from Bioss Antibodies, Woburn, MA, USA; Clarity Western ECL Substrate (1705060) from Bio-Rad, Hercules, CA, USA.

5. Conclusions

Chronic CBD administration (10 mg/kg once a day for two weeks) does not modify BP and HR in a model of primary (SHR) and secondary (DOCA-salt) hypertension and in their respective normotensive controls in spite of the reduction of cardiac and plasma oxidative stress. Whether, besides its direct effect, CBD also possesses an indirect anti-oxidant effect that is based on the endocannabinoid system is questionable. Thus, CBD had opposite effects on numerous components of the endocannabinoid system in both hypertension models. The unexpected CBD-related increase in lipid peroxidation in normotensive controls deserves further investigation and may lead to untoward effects if CBD is used for therapeutical purposes listed in the Introduction. Provided that our data on animals can be transferred to humans, CBD will not lead to an unexpected fall in blood pressure in patients.

Author Contributions: Conceptualization, B.M.; methodology and investigation, P.R., I.J.-K., M.B., A.J., M.T., E.H.-S.; formal analysis, P.R. and A.P.-B.; writing—original draft preparation, review and editing, P.R., E.S., B.M.; visualization, P.R., M.T., A.P.-B., B.M.; supervision, B.M.; project administration, B.M. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by National Science Centre (Poland), grant number 2015/19/B/NZ7/02270. Publication was written during doctoral studies (P.R.) under the project No POWR.03.02.00-00-I051/16 co-funded from European Union funds, PO WER 2014-2020.

Acknowledgments: We wish to thank I. Malinowska and A. Toczydłowska for their excellent technical assistance.

Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

2-AG 2-arachidonoylglycerol
4-HHE 4-hydroxyhexenal
4-HNE 4-hydroxynonenal
AA arachidonic acid
ACN acetonitrile
AEA anandamide
A-1-TG arachidonoyl-1-thio-glycerol
ANOVA analysis of variance
BCA bicinchoninic acid
BP blood pressure
BSA bovine serum albumin
CAT catalase
CBD cannabidiol
CE capillary electrophoresis
CO carbonyl groups
DBP diastolic blood pressure
DEA  docosatetraenoyl ethanolamide
DGLEA  dihomo-γ-linolenoyl ethanolamide
DHA  docosahexaenoic acid
DHEA  docosahexaenoyl ethanolamide
DMF  dimethylformamide
DOCA  deoxycorticosterone acetate or DOCA-salt hypertensive rats
DOCA-salt  DOCA-based method to generate hypertension in rats or DOCA-salt hypertensive rats
DTNB  5,5′-dithiobis-2-dinitrobenzoic acid
ECG  electrocardiography
ECS  endocannabinoid system
EDTA  ethylenediaminetetraacetic acid
ESI  electrospray ionization source
FAAH  fatty acid amide hydrolase
FAME  fatty acid methyl esters
FFA  free fatty acids
FID  flame ionization detector
GC  gas chromatography
GPR  G-protein coupled receptor
GPx  glutathione peroxidase
GSH  glutathione
GSR  glutathione-disulfide reductase
GSSG  glutathione disulfide
HEA  homo-γ-linolenyl ethanolamide
HPLC  high-performance liquid chromatography
HR  heart rate
ID  internal diameter
i.p.  intraperitoneal injection
IS  internal standard
LA  linoleic acid
LEA  linolenoyl ethanolamide
MAGL  monoacylglycerol lipase
MDA  malondialdehyde
MRM  multiple reaction monitoring
MS/MS  tandem mass spectrometry
m-NA  m-nitroaniline
NADA  N-arachidonoyl dopamine
NADPH  nicotinamide adenine dinucleotide phosphate
OEA  oleoyl ethanolamide
O-PFBoxime  O-(2,3,4,5,6-pentafluorobenzyl) oxime
PEA  palmitoyl ethanolamide
PH  phospholipids
PIPS  piperezine-N,N′-bis(2-ethanesulfonic acid)
POEA  palmitoleoyl ethanolamide
PPR  peroxisome proliferator-activated receptor
PUFA  polyunsaturated fatty acids
PVDF  polyvinylidene difluoride
RAAS  renin-angiotensin-aldosterone system
RIPA  radioimmunoprecipitation assay
SBP  systolic blood pressure
SEA  stearoyl ethanolamide
SEM  standard error of the mean
SHAM  sham-operated rats
SHR  spontaneously hypertensive rats
SIM  selected ion monitoring mode
SOD superoxide dismutase
s.c. subcutaneous injection
THC $\Delta^9$-tetrahydrocannabinol
TLC thin-layer chromatography
TNB 5-thio-2-nitrobenzoic acid
TRPV transient receptor potential vanilloid
UPLC ultrahigh performance liquid chromatography
WKY Wistar-Kyoto rats

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