Phthalates Are Metabolised by Primary Thyroid Cell Cultures but Have Limited Influence on Selected Thyroid Cell Functions In Vitro

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Abstract

Phthalates are plasticisers added to a wide variety of products, resulting in measurable exposure of humans. They are suspected to disrupt the thyroid axis as epidemiological studies suggest an influence on the peripheral thyroid hormone concentration. The mechanism is still unknown as only few in vitro studies within this area exist. The aim of the present study was to investigate the influence of three phthalate diesters (di-ethyl phthalate, di-n-butyl phthalate (DnBP), di-(2-ethylhexyl) phthalate (DEHP)) and two monoesters (mono-n-butyl phthalate and mono-(2-ethylhexyl) phthalate (MEHP)) on the differentiated function of primary human thyroid cell cultures. Also, the kinetics of phthalate metabolism were investigated. DEHP and its monoester, MEHP, both had an inhibitory influence on 3’-5’-cyclic adenosine monophosphate secretion from the cells and MEHP also on thyroglobulin (Tg) secretion from the cells. Results of the lactate dehydrogenase-measurements indicated that the MEHP-mediated influence was caused by cell death. No influence on gene expression of thyroid specific genes (Tg, thyroid peroxidase, sodium iodine symporter and thyroid stimulating hormone receptor) by any of the investigated diesters could be demonstrated. All phthalate diesters were metabolised to the respective monoester, however with a fall in efficiency for high concentrations of the larger diesters DnBP and DEHP. In conclusion, human thyroid cells were able to metabolise phthalates but this phthalate-exposure did not appear to substantially influence selected functions of these cells.
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**Introduction**

Phthalates are plasticizers used in a large variety of consumer products and building materials. They are produced as diesters but are rapidly metabolised to monoesters when entering an organism. More complex and larger phthalate molecules are further metabolised to secondary metabolites by oxidation, before they are excreted partly glucuronidated in the urine [1–3]. While the metabolism of phthalates has been previously studied in cell cultures [4–9] the kinetics of phthalate metabolism from di- to monoester are unknown.

It is well known that phthalates have anti-androgenic activity [10;11] and they are also suspected to influence the thyroid axis, reviewed in [12]. Associations between phthalate exposure and thyroid function have thus been investigated in several epidemiological studies [13–20], suggesting that phthalates may influence the concentration of peripheral thyroid hormones [13–15;18;20], though both positive and negative associations have been observed. Similarly, studies in rodents have demonstrated that phthalates may either decrease or increase peripheral thyroid hormone concentrations, with or without a concurrent change in thyroid stimulating hormone (TSH) concentrations [21–26]. One human *in vivo* study failed to detect any influence from dermal phthalate application on circulating TSH or peripheral thyroid hormone concentrations [27], as did one rodent *in vivo* study using oral phthalate exposure [28]. Few *in vitro* studies have investigated phthalate-effects on the thyroid axis and very different endpoints have been used in the study designs [29–36]. Other studies have investigated possible phthalate-mediated effects on the thyroid hormone receptor but are not relevant in relation to this study and therefore not further mentioned here.

The aim of the present study was to investigate if phthalates exerted a direct influence on human thyroid cells in primary cultures. The ability of the human thyroid cells to metabolise phthalates as well as the influence of phthalates on selected functions of these cells were investigated.

**Methods**

**Cell cultures**

Primary human thyroid epithelial cells were cultured as previously described [37] with minor modifications. In brief, paraadenomatous tissue removed during thyroidectomies at the Department of Ear, nose and throat (ENT)-Head and Neck surgery, Rigshospitalet, University of Copenhagen, was washed in phosphate buffered saline (PBS) (calcium and magnesium free, Gibco, Invitrogen Thermo Fischer Scientific, Waltham, MA, USA) and cut into small pieces followed by incubation with collagenase I (Sigma-Aldrich, St. Louis, MO, USA) and dispase II (Roche, Basel, Switzerland) for 75 minutes at 37°C. The digested tissue was filtered through a 100 μm filter (Falcon, BD bioscience, NJ, USA) and HAM’s F-12 culture medium supplemented with L-glutamin (Panum Institute, Copenhagen University, Denmark), 5% foetal bovine serum (FBS) (Biological Industries, Beit HaEmek, Israel), non-essential amino acids, penicillin and streptomycin (Gibco) were added. The suspension was centrifuged at 1200 x G for 5 minutes, and cells were re-suspended in HAM’s F12 culture medium containing the same supplements as mentioned above and six additional nutritional factors: TSH (1U/l, Sigma-Aldrich), insulin (Eli Lilly, Herlev, Denmark), transferrin and glycyll-histidyl-lysine acetat (Sigma-Aldrich), somatostatin and hydrocortisone (Calbiochem, EMD Millipore, Billerica, MA, USA). Cells were seeded on 24 well plates and cultured in humidified air, 37°C, 5% CO₂, to confluent monolayers for approximately 10 days. Before initiation of the experiments, cell cultures were starved from TSH for 3 days. Experiments were conducted using culture media without FBS and in presence or absence of TSH (i.e. TSH-stimulated or unstimulated cultures).
Substances to be tested (phthalates or a positive control, interleukin (IL) -1β) were added separately to experiment wells and 0.1% ethanol was added to the negative controls of phthalate experiments. Cell cultures were incubated for 24, 48 or 72 hours, before cell supernatants were harvested and stored at -20°C. The cells were harvested on ice immediately after by using lysis buffer (Qiagen, Hilden, Germany) and stored at -80°C with prior addition of 70% ethanol.

Experiments were conducted either in single determination, duplicates or triplicates, which is specified in the result section. Replicates derived from the same cell culture were grown on the same 24-well culture plate.

Primary thyroid cell cultures obtained by the described method were shown to contain 98% thyroglobulin (Tg)-producing cells [38]. The function of each thyroid cell culture was ensured by assessing the outcome variables from both TSH- and unstimulated cells on all plates. To optimise the cell cultures, the impact of TSH-starvation, addition of FBS during experiments and the duration of experiments were studied. In these studies, cell supernatants and cells were harvested as described above, except for the study of FBS-addition where no cells were harvested.

Outcome variables in the experiments were Tg and 3’-5’-cyclic adenosine monophosphate (cAMP) secretion as well as gene expression of Tg, thyroid peroxidase (TPO), sodium iodine symporter (NIS), thyroid stimulating hormone receptor (TSHr) and interleukin (IL)-6. These variables were also assessed in the presence of IL-1β (final concentrations of 100, 10³, 10⁴ and 10⁵ U/l and 72 h exposure; Novo Nordisk A/S, Copenhagen, Denmark). The cytokine IL-1β is known to consistently inhibit the differentiated functions of thyroid cells (reviewed in [39]) which in turn produce/secrete cytokines (such as IL-6) instead [40], and IL-1β therefore served as positive control of the experimental design.

Phthalates

Three diesters, di-ethyl phthalate (DEP, CAS: 84-66-2), di-n-butyl phthalate (DnBP, CAS: 84-74-2) and di-(2-ethylhexyl) phthalate (DEHP, CAS: 117-81-7), and two monoesters, mono-(2-ethylhexyl) phthalate (MEHP, CAS: 4376-20-9) and mono-n-butyl phthalate (MnBP, CAS: 131-70-4) were investigated in this study. The function of the thyroid cells exposed to separate phthalates was assessed in both TSH- and unstimulated cell cultures, the latter to investigate if phthalates themselves were able to activate cells. Since pilot studies in unstimulated cultures indicated limited influence by phthalates (data not shown), only DEP, DnBP and MnBP were investigated for Tg and cAMP secretion, and DEP for gene expression in these cultures. On the other hand, TSH-stimulated cultures were exposed to all three diesters (DEHP, DnBP and DEP) and the two monoesters (MEHP and MnBP) with measurement of both Tg- and cAMP-secretion. Furthermore, the influence of the diesters (DEHP, DnBP and DEP) on gene expression was investigated.

DEP and DEHP were purchased from Aldrich Chemical Company (Sigma-Aldrich), DnBP from VWR International (Radnor, PA, USA) or Aldrich Chemical Company, MnBP from Aldrich Chemical Company or Cambridge Isotope Laboratory (Tewksbury, MA, USA) and MEHP from Cambridge Isotope Laboratory. Phthalates were dissolved in ethanol prior to further dilution in the cell culture media (final ethanol concentration was 0.1%) and added separately to the cell culture wells at a final concentration of 0.001, 0.01, 0.1, 1, 10 or 100 μM, respectively. The concentrations of mono-ethyl phthalate (MEP), MnBP and MEHP in cell culture supernatants were quantified by isotope diluted online-Turbo Flow-liquid chromatography-tandem mass spectrometry based on a method previously described [41], but modified to an 11.5 minute runtime. The metabolites were all primary metabolites of DEP, DnBP and DEHP, respectively. Inter- and intra-assay variations of cell media spiked with MEP, MnBP
and MEHP, at four different concentrations showed variations from 2.9% to 8.8%. All results are given as mean ± SD.

To investigate how fast metabolism occurred, a time-study was conducted. Primary thyroid cell cultures were exposed to either DEP, DnBP or DEHP at 0.1 μM for 0, ¼, ½, 1, 2, 4, 24, 48 and 72 hours, after which cell culture supernatants were harvested and their contents of MEP, MnBP and MEHP were determined.

cAMP

3-Isobutyl-1-methylxanthine (IBMX, Sigma-Aldrich) was added to cell cultures used for cAMP assessment concurrently with the phthalates or IL-1β. IBMX was diluted in ethanol (final ethanol concentration 1%), and thus, 1.1% ethanol was added to the ethanol-control in these experiments. Cells were harvested as described above, and the cAMP concentration was measured by a competitive protein binding method [42]. The intra-assay variation at concentrations of 0.4 and 1.4 μM was 4.7 and 7.2%, respectively (n = eight duplicates for each control level). The inter-assay variation was 13.5% for the low control (range 0.29–0.45 μM) and 9.7% for the high control (range 1.10–1.71 μM) (n = five samples in duplicates for each control). The calibration range was 0.05 to 2.0 μM.

Thyroglobulin

Secretion of Tg was measured in supernatants by enzyme-linked immunosorbent assay (ELISA). Wells of polystyrene plates were coated with mouse Tg-antibody (Tg-Ab) (AbD Sero-tec, Oxford, UK). After addition of supernatants, rabbit anti-human Tg-Ab [43] was added, and plates were incubated for 60 minutes at 37°C, washed and peroxidase-conjugated polyclonal swine anti-rabbit immunoglobulin was added together with murine serum (both DAKO, Glostrup, Denmark). After 60 minutes incubation, plates were washed and a chromogenic substrate was added (TMB One, KEM EN TEC diagnostics, Taastrup, Denmark). The reaction was stopped by adding 0.18 M sulphuric acid and results were read using an ELISA reader (BioTek Synergy 2, Winooski, VT, USA) at 450 nm. The intra-assay variation at 52 and 101 μg/l was 9.5 and 8%, respectively (n = seven and six duplicates for the low and high control level, respectively). The inter-assay variation was 22.3% for the low control (range 32–65 μg/L) and 17.5% for the high control (65–121 μg/L) (n = five samples in duplicate for each control). The standard range was 10 to 500 μg/L.

RT-qPCR

Total RNA from harvested primary human thyroid cells was extracted with an Rneasy mini kit (Qiagen) according to the manufacturer’s protocol. However, DNA elimination was not performed routinely. The concentration and purity of the achieved RNA was measured on a NanoDrop spectrophotometer (nd-1000, Wilmington, DE, USA). cDNA was synthesized (Superscript VILO synthesis kit (Invitrogen)) by mixing 4 μL of the VILO reaction mix, 2 μL of the Superscript enzyme mix, the same amount of RNA from each sample and RNAase free water to a total volume of 20 μL. Samples were incubated for 10 minutes at 25°C, 60 minutes at 50°C and 5 minutes at 85°C, after which 80 μL of 0.5X Tris-EDTA-buffer (Sigma-Aldrich) were added.

For real-time quantitative polymerase chain reaction (RT-qPCR), SYBR Green JumpStart Taq Ready Mix (Sigma-Aldrich) was used. Primers and the respective sequences are listed in S1 Table. A pool of undiluted cDNA was used for preparation of the standard curve. Four μL of SYBR Green JumpStart Taq ReadyMix, 10 μL H₂O and 1 μL primer-mix (1μM final concentration of each primer) were added to each reaction. RT-qPCR was done on Lightcycler 480 II
(Roche) with the following cycling: Initial denaturation at 94°C for 2 minutes, followed by 45 cycles of 30 seconds at 94°C and 45 seconds at 59°C and 1.30 minutes at 72°C, and final melting curve analysis. The high number of cycles was used due to very low expression of both NIS and IL-6. The specificity of the amplified products was verified by melting curve analysis and the relative mRNA quantification was achieved by standard curve method. The obtained gene of interest-expression was then divided by the housekeeping gene (HKG)-expression of the same sample.

Four different HKG were used: beta-2-microglobulin (B2M), actin, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) or ATP synthase H+ transporting mitochondrial F1 complex beta polypeptide. Neither phthalates, TSH nor ethanol had an influence on these HKG. At least two HKG were included in the analysis of each experiment and the genes of interest were normalised to the most stable HKG. Stability was judged by the standard deviation of the HKG and by correlation to other HKG from the same experiment.

Cytotoxicity
Phthalate-induced cytotoxicity was evaluated by measurement of lactate dehydrogenase (LDH) content in cell culture supernatants. A homogenous membrane integrity assay (CytoTox-ONE, Promega, Fitchburg, WI, USA) was used according to the manufacturer’s protocol with the modification that the LDH content was assessed in harvested supernatants instead of directly in the cell cultures. Briefly, one thyroid cell culture was exposed to DEHP and MEHP (0.001 to 100 μM), and both a positive Triton X-100 (0.02, 0.2 or 1.9 mg/L) control and three cell culture controls (unstimulated control, TSH-stimulated control and ethanol control) were added. Culture supernatants were harvested after 72 h of incubation and centrifuged at 1200 x G for 5 minutes. The supernatants were then transferred to a black 96-well half area microplate (Th. Geyer, Renningen, Germany) followed by addition of the same volume CytoTox-One reagent to all wells. The microplate was stirred in a shaker and incubated at room temperature for 10 to 15 minutes. Hereafter, stop solution (included in the assay) was added, and the plate was again stirred before results were read on a fluorometer (Victor2, PerkinElmer, Waltham, MA, USA). The LDH contents/amount of lysed cells was proportional to the fluorescence produced and given in relative fluorescence units, as a measure of toxicity of the phthalates.

Ethics
This study was approved by The Danish Committees on Health Research Ethics, Capital Region (Protocol number: H-1-2012-110), which in Denmark also represents the institutional review board. Oral and written informed consent was given by the study participants before donation of the thyroid tissue used for primary thyroid cell cultures.

Statistics
Results from the method evaluation (except evaluation of incubation time) were analysed by paired T-test. For all other experiments, 2 Way ANOVA and Tukey’s Post-hoc analysis were used to compare experiment groups (SAS institute). All analyses were performed with either untransformed or log10-transformed data as appropriate (which is noted in the tables or figure legends), depending on homogeneity of variance (Bland-Altman plot) and normal distribution (histogram). If experiments were conducted in duplicates or triplicates, the mean of the outcome variable was used in analysis. P-values <0.05 were considered statistically significant.
Results

Thyroid cell cultures

Concentrations of cAMP and Tg in supernatants and expression of Tg-, TPO-, NIS- and TSHr-mRNA were significantly lower in unstimulated compared to TSH-stimulated thyroid cells (Table 1). TSH-starved cell cultures had higher cAMP-levels compared to non-starved cells, but only in TSH-stimulated cultures (p = 0.03, n = eight cultures in triplicates). None of the other functional variables were significantly influenced by starvation, neither in TSH- nor unstimulated cultures (S2 Table).

Absence of FBS during experiments resulted in higher cAMP levels and unchanged Tg levels compared to experiments conducted in presence of FBS (p = 0.04, three cultures in triplicates), yet, only observed in TSH-stimulated cultures (S2 Table).

For evaluation of incubation time, cells and supernatants were harvested after 6, 12, 48 or 72 hours, respectively. All assessed variables increased over time. Thus, the Tg-secretion was higher after 72 hours than at all other time points, whereas cAMP-secretion stagnated after 48 hours (S3 Table). Gene expression of all assessed genes was higher after 24, 48 or 72 hours compared to 6 hours. However, there was no significant difference between these three time points (24, 48 and 72 hours), except for Tg- and TPO- mRNA where expression was higher after 48 and 72 hours, respectively, compared to 24 hours (S3 Table).

The positive control cell cultures exposed to IL-1β for 72 hours in TSH-stimulated cultures demonstrated a dose-dependent inhibition of both Tg and cAMP secretion by this cytokine. Likewise, the expression of all thyroid specific genes was impaired in a dose dependent manner, while gene expression of the cytokine IL-6 was enhanced by IL-1β (S1 Fig).

Metabolism of phthalates

The monoesters MnBP and MEHP, when added to cell cultures, were fully recovered at all added concentrations. All three diesters, DEP, DnBP and DEHP, were metabolised to their respective primary monoester, MEP, MnBP and MEHP. No secondary metabolites of DEHP, such as mono-(2-ethyl-5-hydroxyhexyl) phthalate, mono-(2-ethyl-5-oxohexyl) phthalate or mono-(2-ethyl-5-carboxypentyl) phthalate were detected.

Cell cultures exposed to the low diester-concentrations (0.001 up to 0.1 μM) had higher measurable monoester concentrations than the added diester-concentration. Thus, for instance 0.3 μM MnBP was measured in cell cultures exposed to 0.1 μM DnBP (data not shown). For the high diester-concentrations (1 to 100 μM), DEP was almost fully metabolised to the mono-ester MEP while metabolism of DnBP and DEHP to their respective monoester declined with

Table 1. Levels of outcome variables in un- and TSH-stimulated controls. Experiments were conducted in duplicates or triplicates and exposure time was 72 h. Log10-transformed data was used in statistical analysis. cAMP: 3’-5’-cyclic adenosine monophosphate. IL: interleukin. NIS: sodium iodine symporter. Tg: thyroglobulin. TPO: thyroid peroxidase. TSHr: thyroid stimulating hormone receptor.

| Outcome variables | n | unstimulated control | TSH-stimulated control | unit | P-value |
|-------------------|---|----------------------|------------------------|------|---------|
| cAMP              | 23| 0.00002 (0–0.15)     | 0.48 (0.06–4.04)       | μM   | <0.0001 |
| Tg                | 22| 556 (76–72,052)      | 1020 (104–83,649)      | µg/L | 0.003   |
| Tg mRNA           | 17| 0.2 (0.01–1.2)       | 0.7 (0.01–3.5)         | ratio to housekeeping gene | 0.0008 |
| TPO mRNA          | 18| 0.2 (0.04–1.1)       | 1.2 (0.1–4.4)          | ratio to housekeeping gene | <0.0001 |
| NIS mRNA          | 16| 0.04 (0–2.9)         | 1.5 (0.4–72)           | ratio to housekeeping gene | 0.0001 |
| TSHr mRNA         | 18| 0.8 (0.2–2.3)        | 1.0 (0.08–3.4)         | ratio to housekeeping gene | 0.04   |

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increasing concentrations of the diester (Fig 1). There was a time-dependent increase in the monoester content of all three added diesters (0.1 μM). The highest contents of MnBP, MEP and MEHP were found after 24, 48 and 72 hours, respectively. After 4 hours, 86%, 93% and 25% of the respective maximal content of MEP, MnBP and MEHP were observed (Fig 2).
After 72 hours of incubation in TSH-stimulated cultures, both DEHP and MEHP had a significant influence on cAMP-secretion (DEHP: p = 0.0006, n = nine cultures in single determination; MEHP p = 0.01, n = three cultures in single determination) (Fig 3A). Comparison of exposure groups by post-hoc analysis demonstrated an inhibition by 10 μM DEHP compared to the control, 1, 0.1 and 0.01 μM, and an inhibition by 100 μM MEHP compared to all other concentrations (S4 Table). DnBP seemed to stimulate cAMP-secretion, which however was not
statistically significant (Fig 3A and S4 Table). Studying the single courses of these experiments revealed the following: in three out of eight experiments, DnBP stimulated cAMP-secretion, which was very marked for one culture; in the remaining other five cultures the cAMP-secretion was either below or about the same levels as that of the ethanol control (S2 Fig).
None of the investigated phthalates had any influence on cAMP-secretion in unstimulated cultures (S4 Table).

Only MEHP had a significant influence on Tg-secretion from TSH-stimulated cultures after 72 hours of incubation (p = 0.01, n = three cultures in single determination) (Fig 3B). Comparison of exposure groups by post-hoc analysis demonstrated an inhibition by 100 μM MEHP compared to all other concentrations. None of the investigated phthalates had any influence on Tg-secretion in unstimulated cultures (S4 Table).

Gene expression of all investigated genes (Tg, TPO, NIS, TSHr and IL-6) in TSH- and unstimulated cultures was not significantly influenced by any of the investigated phthalates (72 h phthalate-exposure) (S4 Table).

A time study (using incubation times of 24, 48 or 72 h) showed no significant influence on cAMP- or Tg-secretion nor on Tg-mRNA-expression in cell cultures exposed to concentrations from 0.001 to 100 μM DEHP (data not shown).

**Cytotoxicity**

The LDH-content of the three negative culture controls (TSH-stimulated, unstimulated and ethanol control) was lower than the positive Triton X-100 control at the two highest concentrations (0.2 and 1.9 mg/L). Supernatants of cell cultures exposed to MEHP at 100 μM had a higher LDH-content compared to all negative culture controls. At all other concentrations, MEHP and DEHP resulted in an LDH-content lower than or similar to the negative culture controls (S3 Fig).

**Discussion**

In the present *in vitro* study, a limited influence on the function of primary thyroid cell cultures by all the investigated phthalate di- and monoesters was found. Experiments with phthalates were mainly conducted in single determination in order to accommodate all dilutions and controls for comparison on the same culture plate, while the number of experiments was sought to be at least nine. Since no major dose-response effects were detectable, especially compared to the positive control IL-1β, some phthalates and outcomes were studied in less than nine experiments. Thus results are only indicative, but still conclusive in that no major or dose-dependent influences were found.

Only DEHP at the second highest concentration (10 μM) inhibited cAMP-secretion. MEHP at the highest concentration (100 μM) inhibited both cAMP- and Tg-secretion but also resulted in an increased LDH-release. Thus, the observed influences at this high concentration were probably caused by cell death resulting in less production of cAMP and Tg, rather than mere modulation of the cellular function. Also, a high ethanol concentration (1.1%) was present in cultures where cAMP was measured and could itself act in synergy with the phthalates and result in a toxic influence. Only one other *in vitro* study has investigated the influence on cAMP production (human recombinant (hr)-TSHr-mediated) by phthalic acid where no significant influence was described [34].

Gene-expression of Tg, NIS, TPO and TSHr, which increased with culture time and were therefore chosen to be assessed after 72 h of incubation, were unaffected by DEP, DnBP or DEHP-exposure in the present study. A recent *in vivo* study by Liu et al. of Sprague-Dawley rats demonstrated a decrease in serum TPO, NIS and peripheral thyroid hormones, but unchanged concentrations of serum Tg, TSH and thyrotropin releasing hormone by DEHP [44]. The group of Ulrich Loos investigated how phthalates influenced the iodine uptake and NIS gene expression in rodent cell lines (FRTL-5 and PC C13, respectively) [29;36]. DEHP but not di-butyl phthalate (DBP) increased iodine uptake, while neither DEHP nor DBP affected
NIS mRNA expression, which on the other hand was influenced by other investigated phthalates [29;36]. Song et al. investigated the activity of the hrTPO gene and found it to be increased by DBP (100 μM) [35]. In summary, no other or only few published studies have investigated phthalate mediated influences on Tg, NIS, TPO or TSHr. Of these studies, four (including the present study) have investigated NIS. Thus, DEHP did not seem to affect the gene expression of NIS, but may influence iodine uptake. NIS in serum, which was analysed by a commercial ELISA-kit [44], could also derive from other cells than thyroid cells [45], making assumptions on phthalate mediated influence on the thyroid-derived NIS uncertain.

To confirm that our cell system was suitable for investigation of inhibiting substances, IL-1β, was used as a positive control since its influence on the release of Tg and cAMP from primary and secondary human thyroid cell cultures has repeatedly and reproducibly been demonstrated [37;40;46–48]. In the present study, a similar influence on thyroid specific gene expression level was seen. Thus, IL-1β caused a dose-dependent decrease in thyroid specific genes (TPO, TSHr, NIS and Tg) but increased the IL-6 gene expression [40]. Only one experiment with IL-1β was conducted in the present study and judged sufficient since results confirmed those of numerous previous studies with IL-1β in our laboratory [37;40;46–48].

Phthalate-metabolism has previously been studied in cell cultures [4–9]. Kristensen et al. demonstrated an uptake of phthalate diesters but not monoesters by SC5 cells (juvenile mouse Sertoli cell line) and consequent metabolism of the diesters to the respective monoesters [4]. Phthalate metabolites of the present study were assessed in supernatants and thus, no firm assumption can be made about the cellular uptake of the two monoesters, MnBP and MEHP, by primary thyroid cells. However, when diesters were added to thyroid cells, the monoester concentration was increased dose-dependently, suggesting that the diesters were taken up and metabolised by the primary thyroid cell cultures. The extent and timing of metabolism differed for the three diesters: DEP was metabolised at all concentrations, while DnBP and DEHP were metabolised less effectively with increasing concentrations. DnBP and DEHP are less water-soluble than DEP and may not have been dissolved properly at the high concentrations, reducing their availability to the cells. Another observation was that DEP and DnBP appeared to be metabolised faster than DEHP, which could be due to different mechanisms of uptake into the cells due to the larger molecular size of DEHP.

FBS contains proteins, immunoglobulins and enzymes, and could therefore influence the assays used for both Tg and cAMP-quantification and also the metabolism of phthalates. FBS was not present during the experiments but other substances present in the culture media during the experiments could potentially influence the metabolism of phthalates. Phenol red has for instance oestrogen-like activity [49] and oestrogen receptors are also present in normal thyroid tissue [50]. Thus, phenol red may influence the function of thyroid cell cultures and thereby their ability to metabolise phthalates, which needs further investigation.

The phthalate concentration range used in this study included doses likely to be found in humans. Urinary concentrations of MEP in healthy Danish adolescents can reach levels as high as 13 mg/l (67 μM) [51]. In high-exposed groups, such as preterm children, the MnBP concentration in urine reached as high as 6.5 mg/l (29 μM) [52]. Since no previous study has used primary human thyroid cultures, and thus no data exist on the potency of phthalates on the function of these cells, we decided to study single phthalates in a broad concentration range instead of phthalate mixtures, which eventually may be a more realistic in vivo scenario. When phthalates were added to cell cultures at low concentrations (0.001, 0.01 or 0.1 μM), the measured monoester concentrations were higher than expected. In previous studies performed in our laboratory, both high background contamination of some monoesters (probably through phthalate containing laboratory equipment), and migration of some phthalates
to adjacent wells were found [53]. One or both of these observations may account for the unexpected high monoester concentrations found in the present study. Thus, there is a risk that the observed influences on Tg and cAMP secretion in this study may have been caused by a mixture instead of single phthalates, since this could not be corrected for in the statistical analyses.

The purity and correct function of thyroid cells in the cultures have previously been investigated by measuring both the content of Tg-producing cells [38], and the responsiveness to TSH resulting in increased cAMP and Tg secretion, since TSH is the specific stimulator of thyroid cells both in vivo and in vitro [54]. The hormonal end products, triiodothyronine and thyroxine were previously shown to be released from the human thyroid cultures in very small quantities, probably due to absence of iodine in the culture media (own unpublished data). In the present study, the secretion of cAMP, Tg as well as gene-expression of thyroid specific genes (Tg, TPO, NIS and TSHr) were increased in TSH-stimulated cell cultures, confirming their differentiated thyroid cell functions. The 2% non-Tg producing cells present in primary thyroid cell cultures could theoretically be immune cells. As we also assessed the cell cultures gene expression of IL-6, this cytokine mRNA could theoretically originate from these immune cells, but not very likely considering the concentrations.

A limitation of studies with primary thyroid cell cultures in monolayer is their structural difference compared to thyroid cells arranged as follicles in vivo. Monolayers have a different cellular polarity and may also differ in the connections in-between cells compared to in vivo follicles. Consequently, phthalates may influence thyroid cells differently in vivo compared to the in vitro monolayers.

**Conclusion**

Human thyroid cell cultures were able to metabolise phthalates from diesters to the respective monoesters. Most of the metabolism occurred during the first 24 hours and the cells metabolised DnBP and DEP faster than DEHP. A less effective metabolism of high concentrations of DnBP and DEHP was also observed. Only DEHP at 10 μM and MEHP at 100 μM were able to influence Tg- and/or cAMP-secretion. Though only investigated for DEHP, gene expression of thyroid specific genes were unaffected by DEHP exposure. For MEHP, the findings were probably caused by toxicity indicated by an increase in the supernatant-LDH-content. Though some experiments were conducted in low numbers and results thus are only indicative, no sign of major non-toxic or dose-dependent influence were seen in our studies. Proposed hypotheses from epidemiological studies of a direct effect of phthalates on the thyroid gland were not substantiated by the present in vitro study of differentiated gene expression as well as cAMP and protein synthesis from cultures human thyroid cells. An effect of the phthalates on thyroid hormone secretion is therefore unlikely, and such possible effects should therefore be looked for in other parts of the thyroid axis [55].

**Supporting Information**

S1 Fig. Function of TSH-stimulated human thyroid cell cultures exposed to IL-1β. Cell cultures were exposed to IL-1β (100 to 10^5 U/l) for 72 hours (n = one culture in single determination). The dashed line indicates the level of the ethanol controls (ratio = 1). Tg- and cAMP-secretion was assessed in supernatants and mRNA in cells. cAMP: 3'-5'-cyclic adenosine monophosphate. Tg: thyroglobulin. TPO: thyroid peroxidase. NIS: sodium iodine symporter. TSHr: thyroid stimulating hormone receptor. IL: interleukin.
S2 Fig. cAMP secretion from individual cell cultures (culture 1–7) exposed to DnBP (0.001 to 100 μM) for 72 hours. The dashed line indicates the level of the ethanol controls (ratio = 1). DnBP: di-n-butyl phthalate.

S3 Fig. LDH-contents (y-axis) in supernatants from TSH-stimulated thyroid cell cultures. Cells were exposed to DEHP and MEHP, as well as positive cytotoxicity- (i.e. cell cultures exposed to Triton X-100) and negative culture- (i.e. cell cultures not exposed to phthalates) controls for 72 h. N = one culture in single determination except for the negative controls. The LDH-content was proportional to the produced fluorescence (given in relative fluorescence units (RFU)). DEHP: di-2-ethylhexyl phthalate. MEHP: mono-2-ethylhexyl phthalate.

S1 Table. Sequences of the primers used in RT-qPCR.

S2 Table. The impact of TSH-starvation (A) and addition of FBS (B) during experiments. Results of the outcome variables were analysed by paired T-test and experiment duration was 72 h. A: outcome from experiments with thyroid stimulation hormone (TSH) -starvation are compared to those without TSH-starvation. B: outcome from experiments without foetal bovine serum (FBS) are compared to those with FBS. Experiments were conducted in triplicates. cAMP: 3’-5’-cyclic adenosine monophosphate. IL: interleukin. NIS: sodium iodine symporter. Tg: thyroglobulin. TPO: thyroid peroxidase. TSHr: thyroid stimulating hormone receptor.

S3 Table. ANOVA and Tukey post-hoc results from the study of the culture duration. TSH-stimulated primary thyroid cells were cultured for 6, 24, 48 or 72 h, before supernatants and cells were harvested and outcome variables were analysed. All experiments were conducted in triplicates. cAMP: 3’-5’-cyclic adenosine monophosphate. IL: interleukin. NIS: sodium iodine symporter. Tg: thyroglobulin. TPO: thyroid peroxidase. TSHr: thyroid stimulating hormone receptor.

S4 Table. Overview of 2 way ANOVA and post-hoc Tukey results from phthalate-exposed (72 h) thyroid cell cultures, in TSH- and unstimulated controls. If log10 of data was used, the estimated differences between groups are expressed as ratios, e.g. the mean cAMP-secretion from 10 μM DEHP-exposed cells was 21% lower than the mean cAMP-secretion from 0.001 μM DEHP-exposed cells, and lies with 95% certainty between 37% below and 0% below the mean of 0.001 μM DEHP-exposed cells. The hyphen (-) indicates that no post-hoc analysis was made due to insignificant 2 way ANOVA results. cAMP: 3’-5’-cyclic adenosine monophosphate. DEHP: di-2-ethylhexyl phthalate. DEP: di-ethyl phthalate. DnBP: di-n-butyl phthalate. MEHP: mono-2-ethylhexyl phthalate. MnBP: mono-n-butyl phthalate. IL: interleukin. NIS: sodium iodine symporter. Tg: thyroglobulin. TPO: thyroid peroxidase. TSH: thyroid stimulating hormone. TSHr: thyroid stimulating hormone receptor.

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**Author Contributions**

Conceived and designed the experiments: JFH MMB HF EL MLHN ÅKR UFR. Performed the experiments: JFH MMB EL. Analyzed the data: JFH MMB HF EL MLHN. Wrote the paper: JFH MMB MB CHN EL JHB MLHN TF KMM KB ÅKR UFR.

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