Quantitative profiling of endocannabinoids and related N-acylethanolamines in human CSF using nano LC-MS/MS

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Abbreviations:

CSF: cerebrospinal fluid; NAEs: N-acyethanolamines; MRM: multiple reaction monitoring; RSD: relative standard deviation; CB: cannabinoid; FAAH: fatty acid amide hydrolase; MAGL: monoacyl glycerol lipase; AEA: anandamide; 2AG: 2-arachidonoylglycerol; O-AEA  O-arachidonoyl ethanolamine; NADA:  N-arachidonoyldopamine; 2-AGE: 2-arachidonoylglycerol ether; LEA:  N-linoleylethanolamine; PEA:  N-palmitolethanolamine; SEA:  N-stearylethanolamine; OEA:  N-oleylethanolamine; DHEA:  N-docosahexaenylethanolamide; DEA:  N-docosatetraenylethanolamide; DGLEA:  N-dihomo-γ-linolenylethanolamide; TRPV1: transient receptor potential cation channel, subfamily V, member 1; ID: internal diameter; STD: standard solution; ISTDs: internal standards; LLE: liquid-liquid extraction; ACN: acetonitrile; EtOH: ethanol; LOD: limit of detection; LOQ: limit of quantification; QC: quality control
Abstract

Endocannabinoids, a class of lipid messengers, have emerged as crucial regulators of synaptic communication in the central nervous system (CNS). Dysregulation of these compounds has been implicated in many brain disorders. Although some studies have identified and quantified a limited number of target compounds, a method that provides comprehensive quantitative information on endocannabinoids and related N-acylethanolamines (NAEs) in CSF is currently lacking as measurements are challenging due to low concentrations under normal physiological conditions. Here we developed and validated a high-throughput nano liquid chromatography-electrospray ionization mass spectrometry (nano LC-ESI-MS/MS) platform for the simultaneous quantification of endocannabinoids (anandamide (AEA), 2-arachidonoyl glycerol (2-AG)), ten related NAEs and eight additional putatively annotated NAEs in human CSF. Requiring only 200 µL of CSF, our method has limits of detection from 0.28 to 61.2 pM with precisions of RSD <15% for most compounds. We applied our method to CSF from 45 healthy humans and demonstrated potential age and gender effects on concentrations of endocannabinoids and NAEs. Notably, our results show that docosahexaenoyl ethanolamide (DHEA) concentrations increase with age in males. Our method may offer new opportunities to gain insight into regulatory functions of endocannabinoids in the context of (ab)normal brain function.

Keywords: endocannabinoids, brain lipids, quantitation, tandem mass spectrometry, cerebrospinal fluid, liquid chromatography, lipidomics, miniaturization
Introduction

The endocannabinoid system has emerged as a crucial regulator of synaptic communication in the central nervous system (CNS)(1–3). Dysregulation of this system is implicated in various neurological and psychiatric disorders, such as neuroinflammation, stroke, brain trauma, anxiety and depression(4–8). The endocannabinoid system consists of endogenous lipid messengers (endocannabinoids) that activate two distinct cannabinoid (CB₁ and CB₂) receptors and the enzymes responsible for the synthesis and degradation of the endocannabinoids. Anandamide (N-arachidonoyl ethanolamine; AEA) and 2-arachidonoylglycerol (2-AG) are the two best studied endocannabinoids(9). They are synthesized on demand from membrane lipids in a stimulus-dependent manner and are degraded by fatty acid amide hydrolase (FAAH) and monoacyl glycerol lipase (MAGL), respectively(10, 11).

In addition to AEA & 2-AG, there are endogenous lipid compounds capable of activating the cannabinoid CB₁ and CB₂ receptors, i.e. virodhamine (O-arachidonoyl ethanolamine; O-AEA), N-arachidonoyldopamine (NADA) and noladinether (2-arachidonoylglycerol ether; 2-AGE), but their physiological relevance is largely unknown. Related bioactive lipids, such as N-linoleoylethanolamine (LEA), N-palmitoylethanolamine (PEA), N-oleylethanolamine (OEA) and N-stearoylethanolamine (SEA) are produced through the same biosynthetic pathway as anandamide. These N-acylethanolamines (NAEs) have negligible or only weak affinity for the cannabinoid CB₁ and CB₂ receptors, but are able to indirectly modulate cannabinoid receptor activity by interfering with endocannabinoid metabolism, the so-called ‘entourage effect’(12). Of note, some of the NAEs may exert biological activities through other proteins, such as transient receptor potential
cation channel, subfamily V, member 1 (TRPV1), G-coupled protein receptors (GPR55, GPR18 and GPR119) or nuclear receptors (peroxisome proliferator-activated receptor α and γ(13, 14). It is worth mentioning that the endocannabinoids, arachidonoyl glycerol’s and fatty acid ethanolamides, are not only metabolized by FAAH or MAGL but are oxidized by several other enzymes such as cyclo-oxygenase-2 (COX-2), cytochrome P450 and lipoxygenase (LOX) enzymes. These enzymes convert endocannabinoids into a new array of oxygenated intermediates such as prostaglandin ethanolamides (called as prostandides) and prostaglandin glycerol esters or the lipids can be lipooxegenated to many hydroxylated end-products. However, these intermediates are pharmacologically distinct from their parent endocannabinoids and possess different biological functions (15, 16).

Quantification of endocannabinoid levels in the CNS is important to understand their role in human brain disease. Since the cerebrospinal fluid (CSF) is exchanged with brain interstitium, it is regarded as the most relevant accessible body fluid that reflects free endocannabinoid and NAE levels in the CNS (17). However, limited information is currently available on endocannabinoid and NAE levels in the CSF from healthy humans or patients. Previous studies have quantified the endocannabinoids 2-AG and AEA (18–21) and two NAEs (OEA and PEA) (22–24), but not the related lipid messengers.

Under normal physiological conditions, endocannabinoid concentrations are very low (in the nM to fM range) in CSF compared to their levels in blood and tissues. This puts extra demands on the sensitivity of the quantification method. Advanced powerful techniques such as UHPLC (ultra –high performance liquid chromatography) coupled to MS (mass spectrometry) have been employed to quantify endocannabinoids in blood (25–27) and other tissue extracts (28) but these techniques are less suitable for endocannabinoid...
quantification in CSF due to its limited availability. We envisioned that nano LC would offer the opportunity to increase the sensitivity of the analytical system, and a microfluidic chip-based approach would provide the needed sensitivity by allowing injection of a larger sample volume by coupling an on-line enrichment column and small internal diameter (ID) analytical column, resulting in a higher concentration in the eluting peak. In addition, the reduced flow rate of nano LC compared with conventional LC can improve sensitivity through an increase in the ionization efficiency(29). Using microfluidic chip-based technology, incorporating an enrichment column, a six-port valve, a separation column and a nano electrospray tip, allows the coupling of miniaturized separation with a nano electrospray ionization source(30). This design results in lower dead volumes and fewer connections compared to classical nano LC equipment, thereby leading to increased sensitivity and robustness by less peak broadening and dilution effects during transfer of the sample to LC column and later to LC-MS interface(31).

By making use of these advantages, we developed and validated a nano LC-ESI-MS/MS method for the simultaneous quantification of endocannabinoid and related NAEs in clinical (human) CSF. By combining a simple Liquid-Liquid Extraction (LLE) method for sample preparation, a nano LC separation and Multiple Reaction Monitoring (MRM) for detection, we could show that an extended range of endocannabinoids and related molecules can be measured. To show the applicability of this novel method, CSF samples of 45 healthy volunteers were measured to study variation in endocannabinoid levels in relation to gender and age.
Material and methods

Materials

Ultra performance liquid chromatography (UPLC)-grade acetonitrile (ACN) and methanol (MeOH) were purchased from Actu-All Chemicals (Oss, The Netherlands). Formic acid was purchased from Acros Organics (Morris Plains, NJ, USA). Toluene was obtained from Biosolve (Valkenswaard, The Netherlands). EDTA disodium salt was obtained from Sigma-Aldrich Chemie B.V. (Zwijndrecht, The Netherlands). Purified Millipore H₂O was obtained from a Milli-Q PF Plus system (Merck Millipore). NADA, 1-AG, 2-AG, DEA (N-docosatetraenylethanolamide), DHEA (N-docosahexaenylethanolamide), 2-AGE, DGLEA (N-dihomo-γ-Linolenylethanolamide), AEA, SEA, OEA, LEA, PEA, O-AEA, and deuterated standards, NADA-d8, 2-AG-d8, DHEA-d4, AEA-d8, SEA-d3, OEA-d4, LEA-d4, PEA-d4 were purchased from Cayman Chemical (Ann Arbor, MI, USA).

Preparation of standards and internal standards

Pure standards (>98% purity) in ethanol or acetonitrile at different stock concentrations were provided from the manufacturer. The standard stock solutions (STD) were diluted to a concentration of 1 mM using acetonitrile. A final mix of working solution at 100 nM for compounds in group A (NADA, AEA, DEA, DHEA, DGLEA, 2-AGE, LEA, O-AEA) and at 1 mM for compounds in group B (2-AG, 1-AG, SEA, OEA, PEA) were prepared. The deuterated internal standard solutions (ISTDs) were diluted to 100 μM except for 2-AG-d8 at 259 μM. A final working solution was prepared with 100 nM AEA-d8, NADA-d8, DHEA-d4 and 1000 nM 2-AG-d8, SEA-d3, PEA-d4, LEA-d3 and OEA-d4.
Calibration curve preparation

On the day of analysis, the STD working solution was further diluted to seven additional calibration levels as shown in Table 1. Each calibration point was mixed with ISTD-mix at 1:1. The concentrations of ISTDs were chosen to be near the endogenous concentrations of the analytes in CSF.

Sampling protocol for clinical CSF samples, used for method development and validation

CSF was sampled via lumbar puncture between L3/L4, L4/L5, or L5/S1. CSF was collected in 12-mL polystyrene tubes (no. 160172; no. 160172; Greiner Bio-One B.V., Alphen aan den Rijn, The Netherlands) and processed within 30 minutes. A small portion was used for cell count, and the remaining CSF was centrifuged at 21°C for 5 minutes (2000 rpm, 500 g). A portion of the supernatant was used for clinical measurements (i.e. glucose, total protein), and the remaining CSF was stored in 2.0-mL polyethylene cryotubes (Micro tube, no. 72609; Sarstedt AG & C0, Nümbrecht, Germany) and stored at -20 °C. All samples remained at -20 °C until they were pooled in a 50-mL polypropylene tube (no. 210261; Greiner Bio-One B.V). The pooled sample was aliquoted into 2-mL Eppendorf tubes (no. 211-2120; VWR, Amsterdam, The Netherlands) and stored at -80 °C. These clinical CSF samples were used for method validation purpose and termed “pooled CSF samples”. On the day of analysis, 1000 μL of ethanol was added to each aliquot (0.5 mL CSF) to model conditions of study samples and mixed thoroughly and placed on ice.

Sampling of healthy subjects

CSF study samples were collected for an ongoing biochemical study, which was approved by the local medical ethics committee (LUMC). All subjects provided written informed consent before sampling. Prior to CSF
sampling, 6 mL of cold ethanol (ethanol absolute, no. 8098; J.T.Baker, Phillipsburg, NJ, USA) was added to a 15 mL-polypropylene sampling tube (no. 188271; Greiner Bio-One B.V.) and placed on ice. Ethanol was added for denaturation of enzymes present in CSF upon sampling. Subsequently, 3 mL of CSF was sampled directly into the tube and inverted several times to mix CSF and ethanol thoroughly. Immediately after sampling, the CSF/ethanol mixture was divided in 1.5 mL- aliquots into 7 x 1.8 mL-cryotubes (no. 368632; Nunc, Hardenberg, The Netherlands), and placed on dry ice within 30 minutes from sampling. All aliquots were stored at -80°C within 60 minutes from sampling. No extra freeze-thaw cycles were allowed before sample preparation.

**Sample extraction**

Sample extraction was performed on ice. For CSF samples, 600 μL of the CSF/EtOH mixture (200 μL/400 μL) was transferred into 2-mL Eppendorf tubes and 100 μL of 100 mM citrate buffer (pH 5.0) with 25 mM EDTA and 10 μL of deuterated internal standard mixture was added. After adding 550 μL of ice-cold toluene, the tubes were thoroughly mixed for 3 min using a bullet blender (Next Advance, Inc., Averill park, NY, USA), followed by a centrifugation step (4°C / 5000 g / 5 min). Then 475 μL of the upper layer (toluene) was aspirated and transferred into 500 μL Eppendorf tubes. Samples were dried under gentle stream of nitrogen placed on ice followed by reconstitution with 10 μL of water/MeOH (50/50). The reconstituted samples were centrifuged (4°C / 14000 rpm / 5 min) before transferring into vials for LC-MS.

**Chip-based nano LC-MS/MS analysis**

Endocannabinoids and related NAEs were measured using a targeted method on an Agilent 6460 triple quadrupole MS system in conjunction with Agilent 1100/1200 series nano-LC system consisting of a G1375 capillary pump,
G2226 nano pump and a G1376 uWPS autosampler with thermostat (Agilent Technologies, Waldbron, Germany).

Separation was carried out with an Agilent Polaris-HR-Chip 3C18 (G4240-62030, Waldbronn, Germany), which was inserted in the HPLC-Chip Cube MS interface (Agilent Technologies, Waldbron, Germany) operated at ambient temperatures. This chip has a 360-nL enrichment column with a 150 mm (L) x 75 μm (W) separation column and a particle size of 3 μm. The mobile phase was 10 mM formic acid/Water (A) and ACN (B). The analytical gradient was performed in two steps: first, 8 μL of sample was loaded on the enrichment column using the capillary pump delivering a mobile phase in isocratic mode composed of 40% B at a flow rate of 2 μL/min. After the sample was trapped on the enrichment column, it was switched in line with the analytical column. The nano pump gradient started at 45% B and linearly increased up to 80% B in 12 minutes at a flow rate of 400 nL/min. The column was then held at 95% B for 2 minutes before returning to 45% B. The column was re-equilibrated for 5 minutes prior to the next injection giving a total analysis time of 20 minutes. During the analytical gradient, the enrichment column was flushed with 16 μL of 100% B for 8 minutes. At the same time, the injection needle was washed with ACN/H2O (1/1, v/v) using an injection program. The temperature in the autosampler was maintained at 8°C. Electrospray ionization was operated in positive ion mode with a capillary spray voltage set at 1800 V and nitrogen nebulizer gas flow rate of 4 L/min and a temperature of 365°C. For endocannabinoids and related NAEs, individually optimized MRM parameters were determined for each transition for target compounds and ISTDs. The detailed list of MRM parameters and transitions is shown in Table S-1.

To maximize the potential of the platform, MRM transitions for other N-acylethanolamines (NAEs)
were also included. Targets were selected based on: 1) combinations of major fatty acids and ethanolamine (general structure of NAEs) and 2) measurements of CSF by precursor ion scan mode for the characteristic product ion of NAEs, which is the protonated ethanolamine ion at m/z 62 (data not shown). These putatively annotated NAEs are listed as EA (from ethanolamine) followed by molecular mass (m/z) value. Details of MRM transitions included in the method are listed in Table S-1.

Data preprocessing

Peak area integration was performed with Mass Hunter Quantitative analysis (Version B.05.01; Agilent, Santa Clara, CA, USA) and manually inspected. Peak areas of analytes were corrected by the appropriate ISTD and response ratios were calculated and used throughout the analysis.

Method validation

An in-house 3-day validation protocol was used to determine linearity, LOD and LOQ, precision (intra- and inter-batch effect), recovery and matrix effect(32).

Linearity, LOD and LOQ Purchased standards were spiked to both pooled CSF (‘matrix calibration line’) and water as blank matrix (‘academic calibration line’) in seven different concentrations (Table 1) to determine the linearity, limit of detection (LOD) and limit of quantification (LOQ). For method validation, each calibration level was prepared using the sample extraction method explained above four times and injected once from each sample. Responses in the calibration levels were compared by means of t-tests going from low-level to high level. If there was no significance difference between two adjacent levels, then the lower level was excluded from the calibration curve. Regression lines for both matrix and academic line were calculated with 1/X² weighting. If the
differences in slopes of the regression lines between matrix and academic lines were less than 10%, LOD and LOQ values were calculated from academic lines. When the slopes differed, the matrix line was used for these calculations. Each LOD and LOQ was calculated by $3.3 \times \frac{\text{standard deviation of } y\text{-estimate in the regression line}}{\text{slope}}$, and $3 \times \text{LOD}$, respectively.

**Precision** The intra-batch and inter-batch effects were evaluated using the quadruplicate analysis of the pooled CSF samples spiked at three different concentrations (lower level (C2), medium (C4), and high level (C6)) over the three different days of the validation protocol. The pooled CSF was spiked with ISTDs, which behave like the endogenous compounds and are therefore perfectly suited to determine analytical characteristics. Precision was calculated as the relative standard deviation (RSD) of the ratio of target to ISTD. An RSD less than 15% for medium and high levels and less than 20% for the low level was considered acceptable.

**Recovery and matrix effect** Recovery and matrix effects were evaluated using ISTDs spiked to the pooled CSF samples. Recovery was assessed by determining the ratio of the peak areas obtained by spiking ISTDs to the sample before and after extraction. The matrix effect was assessed by determining the ratio of the peak areas obtained by spiking ISTDs to pooled CSF and to water after extraction.

**Biological application in CSF of healthy subjects**

A small study was performed to understand the changes in endocannabinoid levels in the CSF of healthy subjects due to gender and age. The experimental design included 45 individual study samples plus performance indicators. Duplicate samples were prepared and analyzed for 5 samples (15% of total samples). Additionally, 20 quality
control (QC) samples were prepared by pooling individual study samples. A calibration curve was prepared by spiking the calibrations standards along with internal standards to the QC samples. All samples were randomized and each batch includes calibration samples and an even distribution of duplicates, QC samples and blanks.

Data pre-treatment

To detect outliers among QC samples, a principle component analysis (PCA) was used and QCs outside the 95% confidence region were detected and removed as statistical outliers: Two QC samples showed high concentrations for LEA, OEA, PEA and SEA and were detected as statistical outliers. These concentrations were possibly caused by (an) exogenous source(s)(33). After removal of the outliers, the final QC set was used to determine optimal ISTDs for compounds without an isotopically labeled equivalent as an ISTD, as previously described(34). Values from QC samples were used for monitoring data quality and correcting for shifts in MS sensitivity over time(34). After correction with QC samples, relative standard deviations (RSD) of study QC samples were calculated for all compounds (Table 5). Next, corrected data of biological samples (identified compounds only) were analyzed with PCA to detect outliers. In the current data set, one outlier was detected and was removed from further statistical analysis. Finally, RSDs of duplicates were calculated (Table 5) and afterwards one of two samples was randomly selected from remaining duplicate pairs to analyze variability. One duplicate pair was identified as an outlier due to contamination and was excluded.

Statistical analysis

Prior to statistical analysis, study QC corrected data were log-transformed for data normalization. To study whether endocannabinoid and related NAEs concentrations depend on gender or age, multiple linear regression
analysis was applied. For each compound, its concentration was set as the dependent variable with gender and age as predictors. To study whether potential age effects are gender-specific, linear regression was repeated after splitting data by gender. Statistical analysis was performed with R software (version 3.0.3, function \texttt{lm}). All reported p-values are without multiple testing corrections. P-values <0.05 were considered significant.
Result & Discussion

Method development

Although some analytical methods for profiling endocannabinoids and related NAEs in human plasma have been reported (25, 27), methods for CSF are limited to anandamide, 2-AG, PEA and OEA (22, 23). To achieve a high-throughput and sensitive method for detection and quantification of these compounds, two important parameters were optimized. First, liquid chromatography conditions, including gradient and solvent composition, were optimized to obtain an adequate separation of molecules, especially isomers which are indistinguishable by fragmentation pattern, while at the same time maintaining a short chromatographic run time (Figure 1). For example, DGLEA and EA350 are structural isomers, but are chromatographically separated (Figure 3). Second, optimal transitions were determined for each compound to maximize sensitivity, and for each MS/MS transition it was confirmed with standards that there is no cross-talk between compounds. This method uses dynamic MRM, which calculates optimal MRM parameters for multiple analytes using user inputs of retention times windows (delta RT) and cycle time and these parameters remain constant throughout all runs. The individual MRM dwell time is adjusted to maintain constant cycle time which is selected to provide adequate sampling speed for quantitation of narrow LC peaks. When compared to traditional time segmented methods, dynamic MRM methods include fewer ion transitions per unit time during a typical MS scan, which results in longer dwell times for individual ion transitions. The MRM was operated in positive mode due to their higher ionization efficiencies.

For selective detection and quantification of co-eluting metabolites, specific precursor and fragment ions were used for their separation. The fragmentation pattern for N-acylethanolamine derivatives included the characteristic
loss of m/z 62.1 resulting from the fragmentation of the amide bond (Supplementary Figure S3-a). Our chip-based 
nano LC-MS/MS method allowed quantification of 12 endocannabinoids and related NAEs, along with eight 
putatively identified NAEs in CSF (Supplemental Table S1).

The isomerisation from 2-AG to 1-AG due to acyl migration complicates the measurements of the two 
regio-isomers. As shown in Table 2, the sum of 2-AG and 1-AG showed a superior RSD (5-11%) during batch 
measurement compared with 2-AG only (3-38%), which indicated that acyl migration might have occurred during 
sample preparation and measurement. Another contribution to that error could be problems with peak integration, 
as the two isomers are not separated at baseline. Based on this result, the sum of 1-AG and 2-AG was used for 
进一步 data analysis in this paper. In LLE, toluene was previously reported to suppress acyl migration during the 
evaporation step to a large extent(35). However, moderate levels of acyl migration were observed in the current 
study and this is likely due to difference in matrices and sample handling procedures compared to previous studies. 
Moreover, this is the first study to use toluene extraction method for CSF whereas the previous studies were 
demonstrated in plasma (25, 35). Furthermore, it has been reported that elevated temperatures, presence of serum 
albumin in the sample, and high pH values accelerate acyl migration(35, 36). Interestingly, a recent study reports 
that in plasma 1-AG is not an endogenous compound but rather the result of chemical isomerization during 
sample storage, and therefore summing the concentrations of both the isomers will provide meaningful data for 
biological interpretation(27).

It should be noted that non-negligible amounts of SEA and, to a lesser extent, other NAEs (PEA, SEA, 
OEA) were observed in the ethanol which was used during sample collection. This indicates that ethanol used in
the sample collection protocol is a source of contaminants. We also observed contamination of NAEs, especially SEA, OEA, PEA at low levels when using methanol instead of acetonitrile (data not shown). However, only low levels of SEA compared to the endogenous concentration in CSF were seen in the blank (MeOH/H2O) showing the soundness of the analytical method (Supplemental Figure S1). The consequence of the contaminants present in ethanol, from solvents and other potential sources for PEA, SEA, and OEA, was a decline of analytical characteristics (linearity, LOD, LOQ, precision, recovery and matrix effect). Previous studies identified basal concentrations of these compounds in organic solvents, plastics and glassware(33). A recommendation to minimize contamination is to avoid addition of ethanol during CSF sampling or to check for contamination prior to use.

The stability of endocannabinoids and NAEs and their ex-vivo generation, i.e. during and after sampling needs greater attention to avoid variability and inaccuracy of the measurements. To date, the stability of endocannabinoids in CSF was not reported, but in blood NAEs concentrations dramatically increase when withdrawn blood samples are improperly stored. 2AG, 1AG levels were shown to be stable for 2h but a gradual reduction of 2AG was detected after 4h(25). Here, the pooled CSF used for validation was not stored immediately at cold conditions after withdrawal, which might cause increased NAE and 2AG-1AG levels. This has no negative consequences for this study because these pooled samples were only used to assess the validation parameters by spiking the internal standards, and no biological conclusions were drawn from this pooled sample. The study samples, on the other hand, were processed and stored within 30min after sample collection to improve the stability and accuracy of the measurements.
Method validation

Validation was performed using pooled CSF samples. Linearity and sensitivity (LOD and LOQ) are summarized in Table 3. Precision (Intra and Inter-day) values are summarized in Table 4. Recovery and matrix effect are summarized in Figure 2.

Linearity and sensitivity Pooled CSF samples were spiked with 7 levels of calibration standards (Table 1) to determine linearity, LOD, and LOQ for each compound (9 targets). The goodness of fit $R^2$ ranges from 0.995 to 0.999 for all compounds with the exception of SEA ($R^2$ 0.976) confirming the method is linear in the selected calibration range (Supplemental Figure S4). The LOQs are between 0.9 and 61.2 pM for most of the compounds except for those with a contamination problem (PEA and SEA), which shows that the method is suitable for measuring endogenous levels in CSF (Table 5).

Precision The intra- and inter-batch variability were assessed using the standards spiked at three different concentrations (Low (C2), medium (C4) and high (C6)) in pooled CSF samples (Table 4). Quadruplicate sample preparations were performed in each batch, and a total of three batches were processed. The intra-batch RSDs ranged from 1 to 23%. Possible batch-to-batch effects were calculated using the three calibration levels (C2, C4, and C6) measured on three different days. Inter-batch effects ranged from 3 to 21%.

Recovery, matrix effect Recovery and Matrix effect were evaluated using deuterated compounds otherwise used as ISTDs (Figure 2). Recovery was calculated as the ratio of the response of the ISTD of samples spiked before extraction versus samples spiked after extraction. Recoveries ranged from 38 to 57%. These low recoveries are in part due to incomplete recovery of the organic phase during sample extraction. However, these losses are largely
corrected by the use of deuterated ISTDs; when using one of the deuterated ISTDs as internal standard (DHEA-d4) for the other deuterated compounds, recoveries were 81 to 119%. Matrix effect (ion suppression) was calculated as the ratio of the response of the ISTD spiked after extraction in matrix versus water. The matrix effect ranged from 38 to 81%. The ion suppression for these compounds most probably be due to co-eluting phospholipids. SEA-d3 was removed from Figure 2 due to ion suppression by the presence of a high concentration of unlabeled SEA contamination in the sample collection solvents (see above). The compounds have good recovery after correction by an internal standard added prior to extraction. The matrix effects were different between compounds, these values could be considered as acceptable as long as deuterated ISTDs used are able to correct for these factors, or, if no isotopically form of an analyte is available, the matrix effect is reproducible. For target compounds with matching internal standards, the ISTD corrected concentrations were shown to have acceptable reproducibility. Among the detected compounds in the current method, seven compounds have their respective deuterated ISTD for analytical corrections, but for DGLEA and DEA, deuterated standards were not available. For these compounds, LEA-d4 and AEA-d4 respectively were chosen as their ISTDs due to their chemical similarly and their similar elution times.

**CSF endocannabinoids and related NAEs in healthy subjects**

To demonstrate the applicability of the method, we measured 45 human CSF samples from 22 healthy females (mean age 30.9 years ± 12.8 years SD) and 23 healthy males (mean age 39.3 ± 16.5 SD). Nine compounds were detected and quantified (Figure 3).

Endogenous concentrations and ranges of endocannabinoids measured in healthy subjects are listed in
Table 5. The concentrations of AEA and 2-AG are in line with the previous studies (37). In addition to the compounds shown in Table 3, several putatively annotated NAEs were observed in human CSF (Supplemental Figure S2). Of note, an interesting observation in the current CSF samples is the ratio of DHEA/AEA being around 10, which is contrary to plasma, where the ratio is almost equal to 1 (27). This could be that the availability of DHA (Docosahexaenoic acid) is more in CSF than AA (Arachidonic acid), precursors for DHEA and AEA, which are then processed downstream at equal rates. If not, this could indicate the synthesizing enzymes have a preference for DHA to generate DHEA, or alternatively it could be that AEA being a better substrate for FAAH than DHEA. However, further investigation is required to elucidate the exact reason, but this was not the main aim of this study.

The identity of one of the putatively annotated NAE, EA350 peak was assigned as 5(Z),8(Z),11(Z)-eicosatrienoic acid ethanolamide using a commercial available authentic standard and comparing m/z, retention time and fragmentation pattern. The MS/MS fragmentation is shown in supplementary figure S3-b. In similar manner, the identities of all the remaining putatively annotated NAEs should be confirmed by purchasing authentic standards, however these standards are still not commercially available. Although further work needs to be done for annotating all metabolites and understanding the biological implications, this highly sensitive method clearly shows the applicability in measuring a broad range of endocannabinoids and related NAEs.

Although we could not confirm the identity for most of the putatively identified compounds and observed some matrix effect due to ion suppression, we believe that our method with this broader coverage of endocannabinoids and their congeners will help us to understand the biological insights of the endocannabinoid system as a “whole”
in future studies, and to possibly understand the biological role of these putatively identified compounds.

Moreover, this method depicted acceptable reproducibility and sensitivity of these molecules in the application to clinical samples such as the migraine samples.

**Gender and age differences in healthy controls**

It is important for designing large case-control studies to know whether endocannabinoids and related NAEs concentrations show a gender or age dependence. In our study, these factors were analyzed using data from healthy subjects using multiple linear regression analysis. Significant effects of age were present for DHEA. DHEA concentrations were higher in older males (Figure 4). In blood, higher concentrations of docosahexaenoic acid (DHA), the fatty acid of DHEA, were reported in elderly (≥65 years, gender not reported)(38). Intake of DHA can alter DHEA concentrations in blood and brain(39). Additionally, age-related changes in DHA metabolism could play important roles as well(40). There were no significant gender differences in endocannabinoids or related NAEs when corrected for the age imbalance (Supplemental Table S2). This study proves the importance of proper age and sex matching of cases and controls to obtain significant results in studies directed at discovery of endocannabinoid related biomarkers and/or to obtain novel insights in health and disease.
Conclusion

A nano LC-ESI-MS/MS method for the simultaneous quantification of classical endocannabinoids and the related NAEs was developed and validated in human (clinical) CSF. We measured AEA, 2-AG, ten annotated NAEs and eight putatively annotated NAEs, whereas previously only four endocannabinoids had been measured in CSF(22, 23). Requiring only 200 μL of CSF per sample, this sensitive method has LODs reaching pM-fM levels while maintaining good linearity and precision. Application of this method in CSF from healthy humans resulted in novel insights on gender and age effects, i.e. concentrations of DHEA increased with age in males, illustrating the possibilities of this method to study the regulatory functions of endocannabinoids in the context of brain (dys)function.
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References


Footnote

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Tables

Table 1: Stock concentrations of endocannabinoids and related NAEs standards in the calibration line

<table>
<thead>
<tr>
<th>Compound group</th>
<th>Concentrations (nM)a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C0</td>
</tr>
<tr>
<td>A</td>
<td>0</td>
</tr>
<tr>
<td>B</td>
<td>0</td>
</tr>
</tbody>
</table>

aSeven levels of standard solutions (C1 to C7) are spiked to build the calibration line. These standards will be diluted 21 times in CSF matrix for their final concentrations.
Table 2: Difference in %RSD (Relative Standard Deviation) expressed in percentage between 2-AG alone and combined 2-AG and 1-AG

<table>
<thead>
<tr>
<th>Calibration levels</th>
<th>RSD (%)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2-AG</td>
<td>1-AG +</td>
</tr>
<tr>
<td></td>
<td>2-AG</td>
<td>2-AG</td>
</tr>
<tr>
<td>C0</td>
<td>27</td>
<td>5</td>
</tr>
<tr>
<td>C1</td>
<td>38</td>
<td>8</td>
</tr>
<tr>
<td>C2</td>
<td>20</td>
<td>3</td>
</tr>
<tr>
<td>C3</td>
<td>15</td>
<td>11</td>
</tr>
<tr>
<td>C4</td>
<td>19</td>
<td>6</td>
</tr>
<tr>
<td>C5</td>
<td>16</td>
<td>8</td>
</tr>
<tr>
<td>C6</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>C7</td>
<td>3</td>
<td>4</td>
</tr>
</tbody>
</table>
Table 3: Validation parameters: Calibration range, Retention time, LOD (limits of detection), LOQ (limits of quantification)

<table>
<thead>
<tr>
<th>Compound (abbr.)</th>
<th>C</th>
<th>C=C</th>
<th>m/z</th>
<th>Calibration ranges (pM)</th>
<th>Retention time (min)</th>
<th>R²</th>
<th>LOD (pM)</th>
<th>LOQ (pM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEA</td>
<td>16</td>
<td>0</td>
<td>299.5</td>
<td>4.7-4761</td>
<td>13.29</td>
<td>0.96</td>
<td>35.4</td>
<td>107.4</td>
</tr>
<tr>
<td>LEA</td>
<td>18</td>
<td>2</td>
<td>323.5</td>
<td>0.47-476</td>
<td>11.99</td>
<td>0.96</td>
<td>3.2</td>
<td>9.6</td>
</tr>
<tr>
<td>OEA</td>
<td>18</td>
<td>1</td>
<td>325.5</td>
<td>4.7-4761</td>
<td>13.83</td>
<td>0.97</td>
<td>14.4</td>
<td>43.7</td>
</tr>
<tr>
<td>SEA</td>
<td>18</td>
<td>0</td>
<td>327.6</td>
<td>4.7-4761</td>
<td>16.07</td>
<td>0.97</td>
<td>146.9</td>
<td>445</td>
</tr>
<tr>
<td>AEA</td>
<td>20</td>
<td>4</td>
<td>347.5</td>
<td>0.47-476</td>
<td>12.00</td>
<td>0.97</td>
<td>0.28</td>
<td>0.86</td>
</tr>
<tr>
<td>DGLEA</td>
<td>20</td>
<td>3</td>
<td>349.6</td>
<td>0.47-476</td>
<td>12.89</td>
<td>0.96</td>
<td>0.53</td>
<td>1.6</td>
</tr>
<tr>
<td>DHEA</td>
<td>22</td>
<td>6</td>
<td>371.6</td>
<td>0.47-476</td>
<td>11.84</td>
<td>0.96</td>
<td>0.9</td>
<td>2.8</td>
</tr>
<tr>
<td>DEA</td>
<td>22</td>
<td>4</td>
<td>375.6</td>
<td>0.47-476</td>
<td>13.88</td>
<td>0.95</td>
<td>0.31</td>
<td>0.94</td>
</tr>
<tr>
<td>1-AG+2-AG</td>
<td>20</td>
<td>4</td>
<td>378.6</td>
<td>4.7-4761</td>
<td>13.44(1-AG)</td>
<td>0.97</td>
<td>20.2</td>
<td>61.2</td>
</tr>
</tbody>
</table>

“C” and “C=C” = number of carbon atoms and number of double bonds in the fatty acid chain of the molecule, respectively; m/z = mass-to-charge ratio; R² = correlation coefficient; LOD = limit of detection; LOQ = limit of quantification. The LOD and LOQ values of SEA are high due to the contaminant peak present in ethanol used during sample collection.
Table 4: Analysis of pooled CSF samples showing precision (Intra- and inter-batch effect)

<table>
<thead>
<tr>
<th>Compound</th>
<th>Intra-batch precision (%)</th>
<th>Inter-batch precision (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low (C2)</td>
<td>Middle (C4)</td>
</tr>
<tr>
<td>PEA</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>LEA</td>
<td>23</td>
<td>12</td>
</tr>
<tr>
<td>OEA</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>SEA</td>
<td>14</td>
<td>7</td>
</tr>
<tr>
<td>AEA</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>DGLEA</td>
<td>14</td>
<td>5</td>
</tr>
<tr>
<td>DHEA</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td>DEA</td>
<td>9</td>
<td>3</td>
</tr>
<tr>
<td>1-AG+2-AG</td>
<td>11</td>
<td>5</td>
</tr>
</tbody>
</table>

The precision values are expressed in RSD (relative standard deviation). Low, Middle, High are three different concentration (for concentrations see Table1) used to show intra- and inter-batch precision.
Table 5: Concentrations of endocannabinoids and related NAEs quantified in 45 CSF samples of healthy subjects

<table>
<thead>
<tr>
<th>Compound</th>
<th>Concentration (pM)</th>
<th>QC_ RSD (%) (n = 18)</th>
<th>Duplicate_ RSD (%) (n = 4 pairs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Min</td>
<td>Max</td>
<td>Median</td>
</tr>
<tr>
<td>PEA</td>
<td>41.4</td>
<td>270.4</td>
<td>83.7</td>
</tr>
<tr>
<td>LEA</td>
<td>3.6</td>
<td>28.8</td>
<td>9.6</td>
</tr>
<tr>
<td>OEA</td>
<td>22.0</td>
<td>116.6</td>
<td>44.4</td>
</tr>
<tr>
<td>SEA</td>
<td>67.0</td>
<td>310.3</td>
<td>108.4</td>
</tr>
<tr>
<td>AEA</td>
<td>0.5</td>
<td>2.7</td>
<td>1.6</td>
</tr>
<tr>
<td>DGLEA</td>
<td>0.4</td>
<td>2.2</td>
<td>1.0</td>
</tr>
<tr>
<td>DHEA</td>
<td>3.61</td>
<td>24.89</td>
<td>8.48</td>
</tr>
<tr>
<td>DEA</td>
<td>1.2</td>
<td>5.6</td>
<td>2.9</td>
</tr>
<tr>
<td>1-AG +2-AG</td>
<td>36.5</td>
<td>235.3</td>
<td>82.6</td>
</tr>
</tbody>
</table>

The RSD (relative standard deviation) expressed in percentage of QC (quality controls) and duplicates used during the analysis of healthy subjects. SD: Standard deviation
Figures & Figure legends:

Figure 1: Overlay of MS/MS chromatograms of endocannabinoids and related NAEs in standard mix
Figure 2: Recovery and matrix effect of deuterated internal standards in CSF

Recovery and matrix effect values are expressed in percentages. Recovery: Higher values indicate better recoveries.

Matrix effect: Values above 100% implies ion enhancement and below 100% implies ion suppression.
Figure 3: Endogenous peaks of endocannabinoids and related N-acylethanolamines in CSF

X-axis is retention time and Y-axis is response signal
Figure 4: Relationships between DHEA concentrations and age

Linear regression lines are plotted separately for males (straight line) and females (dotted line). Solid circles represent males and solid triangles represent females. Y-axis shows relative concentrations (pM) before log transformation.