Design of an ecological momentary assessment study of exposure to radiofrequency electromagnetic fields and non-specific physical symptoms

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ABSTRACT

Introduction: Idiopathic Environmental Intolerance (IEI) attributed to electromagnetic fields (EMF) refers to self-reported sensitivity mainly characterised by the attribution of non-specific physical symptoms to low-level EMF exposure emitted from sources such as mobile phones. Scientific studies have not provided evidence for the existence of IEI-EMF, but these studies did not resemble the real-life situation or suffered from poor exposure characterisation and biased recall of health symptoms. To improve existing methods for the study of IEI-EMF, an Ecological Momentary Assessment (EMA) study is designed.

Methods and analysis: The study is an EMA study in which respondents carry personal exposure metres (exposimeters) that measure radiofrequency (RF) EMF, with frequent assessment of health symptoms and perceived EMF exposure through electronic diary registration during five consecutive days. Participants will be a selection from an epidemiological study who report to be sensitive to RF EMF. The exposimeters measure electric field strength in 12 frequency bands. Diary questions include the occurrence and severity of 10 non-specific physical symptoms, mood states and perceived exposure to (sources of) EMF. The relationship of actual and perceived EMF exposure and mood with non-specific physical symptoms will be analysed using multilevel regression analysis with time-shift models.

Discussion: The study has several advantages over previous studies, including assessment of personal EMF exposure and non-specific physical symptoms by an ecological method with a minimised chance of recall bias. The within-person design reduces confounding by time-stable factors (eg, personal characteristics). In the conduct of the study and the analysis and interpretation of its outcomes, some methodological issues including a high participant burden, reactivity, compliance to the study protocol and the potential of chance findings due to multiple statistical testing will be accounted for and limited as much as possible.

INTRODUCTION

Some people experience subjective health symptoms in the proximity of (specific sources or frequencies of) radiofrequency (RF) electromagnetic fields (EMF). Sources of RF EMF in the home environment include mobile or digital-enhanced cordless telecommunications (DECT) phones and their base stations, WiFi, microwave ovens, television and radio transmitters. Although there is lack of a validated case definition, when an individual attributes his/her health symptoms to EMF exposure, this is mostly referred to as electromagnetic hypersensitivity. Owing to similarities with other (unproven) environmental intolerances, such as multiple chemical sensitivities, and because scientific evidence of a causal relationship between EMF exposure and symptoms is lacking, the WHO introduced the
broader term Idiopathic Environmental Intolerance (IEI). When afflicted persons attribute their illness to EMF, it is referred to as IEI-EMF. The health symptoms of IEI-EMF are non-specific and differ from person to person. Frequently mentioned symptoms include fatigue, headaches, concentration problems, nervousness and tinnitus. IEI-EMF has been found to be associated with limitations in social and occupational functioning. The prevalence of IEI-EMF in the population is estimated to be 1.5–5%, but a prevalence as high as 13% has also been reported. These differences are due to the population under study (Western countries vs Taiwan), and probably also the instruments or definitions that were used. For the Netherlands, an estimate of the prevalence of IEI-EMF is not yet available.

Scientific studies have not provided convincing evidence for the existence of a causal bioelectromagnetic mechanism for non-specific health symptoms. Alternative explanations for IEI highlight the role of psychological mechanisms, such as hyper vigilance to threat stimuli, attention bias and somatosensory amplification. For IEI-EMF, some findings suggest that nocebo responses account for the symptoms, in which concerns about a perceived harm precede the development of symptoms. Indeed, persons who suffer from IEI-EMF have relatively high levels of mental distress, anxiety, depression and worries about modern life. Several studies have demonstrated a relationship between negative affect and non-specific health symptoms.

Most evidence for the lack of an association between EMF exposure and non-specific physical symptoms is derived from short-term provocation studies in the laboratory, which have been criticised because of their lack of internal and external (ecological) validity. Criticisms include that a visit to the laboratory may cause anxiety that influences the results, that EMF exposure in the experimental setting does not resemble real-life EMF exposure, and that follow-up times are insufficiently long to capture participants’ responses. Observational studies are subject to other forms of bias due to errors in the recall of symptoms (recall bias) and in the assessment of EMF exposure.

The limitations aforementioned can be solved by ecological momentary assessment (EMA). With EMA, the assessment is momentary, on the spot in real life, and captures life as it is lived. More precise and ecologically valid EMA measurement of personal RF EMF exposure can be performed with exposimeters. This produces more valid estimates than other methods, such as self-reported exposure, geo-coded distance from sources of RF EMF (e.g., base stations) or spot measurements. Recall bias of symptoms can be minimised by using EMA diary methods with short-time frames instead of asking participants to retrospectively report (the usual frequency of) symptoms over a prolonged period.

This article describes the design of an EMA study on the relationship between real-life measured and perceived exposure to RF EMF and the real-time experience of non-specific physical symptoms and mood in self-declared electrohypersensitive people. The study intends to minimise sources of bias by using exposimeters to estimate RF EMF exposure and real time on the spot assessment of symptoms.

Objectives
The key objective of the study is to determine whether in a period of a few days non-specific physical symptoms in persons who report to be sensitive to RF EMF can be explained by objectively measured exposure to RF EMF, or by psychological measures such as perceived exposure and mood. Secondary objectives are to study the manifestation of non-specific symptoms in terms of severity and duration of symptoms, the lag time—in hours to days—between exposure and the presentation of symptoms and to characterise RF EMF exposure of persons with IEI-EMF.

METHODS AND ANALYSIS
Study design
Epidemiological panel studies, which have similarities to EMA studies, have been described as “prospective studies that follow a usually small group of individuals intensively over a short time period […] with the objective to study short-term effects of a time-varying environmental exposure.” A main advantage of a panel design is the availability of measurements of exposure and health outcomes at an individual level. The current study is an EMA study in which, for five consecutive days, participants carry a measurement set consisting of an RF EMF personal exposure metre, a so-called exposimeter, a global positioning system (GPS) logger and an electronic diary. The electronic diary assesses health complaints, perceived exposure and mood. It has to be completed directly at frequent, random alarm cues, as this prevents both recall bias and possibly planned high exposures shortly before the time when filling the diary. This design allows for studying whether non-specific physical symptoms are preceded by exposure to EMF, using various latency times, and/or whether these symptoms are related to psychological variables such as perceived exposure and mood.

Selection of study population
Participants will be recruited from respondents of an existing epidemiological study (EMPHASIS) on non-specific physical symptoms and their relation with model-estimated actual and perceived EMF exposure. This study included 6304 persons who were selected from 21 general practices throughout the Netherlands, varying in level of urbanisation and stratified according to the distance of their residences to a mobile telephone base station. The response rate to the written questionnaire was approximately 50%.

Participants will be selected from the respondents to this survey based on self-reported sensitivity to EMF.
measured with a five-point scale. People who indicated to fully or partly agree with the statement “I am sensitive to antennas and devices using wireless communication (eg, for radio, television, mobile phones, wireless internet, etc)” and who gave their consent to use their address to reapproach will be invited by post to participate in the study. All study materials (diaries, exposimeters and instructions) will be delivered at the participants’ homes, where they will be orally instructed about the study procedures.

**Electronic diaries**

Diary methods are considered suitable to examine self-reported events and experiences in their natural, spontaneous context. Benefits of diary methods are that bias in the recall of events and experiences is reduced because the time between the occurrence of an event or experience and its reporting is minimised. Diary methods can appropriately address the research question of what the correlates and antecedents are of within-person variability in daily experiences.

The current study will use electronic diaries. Advantages of electronic diaries include a higher participant compliance than paper and pencil diaries, control of alarm cues and a detailed log file for compliance check.

For diary keeping we will use LG P-500 Optimus One smartphones running on Android 2.3. Because the study population will include persons with IEI-EMF, the phone operates in flight mode without a SIM card. A check of exposure to extremely low frequencies (Emdex Lite, Enertech Consultants, California, USA) and RF (EM Spy 121, Satimo, France) confirmed that the exposure from the smartphone was negligible, that is, below the detection limits.

Special software for the diaries was developed using Java (Android V2.2 or higher). The diary programme is based on software written for Palm-OS personal digital assistants, which has been developed and used by Houtveen et al. A sampling protocol with a mean interval of 2.5 h and random variation of ±30 min will be used that continues from awakening till bedtime. This sampling scheme leads to approximately 8 alarms per day (based on a 16 h awakening period). Diary prompting will only be disabled during sleep, initiated by a button on the smartphone. The smartphone can be used as a morning alarm, and prompting continues after awakening. All unused buttons are blocked. Alarms without response are repeated (maximum 3 times with 10 min time intervals). The alarm software generates a log-file containing alarm and response times to be used for determination of the compliance. The questionnaire can be launched by a start button that is visible for 5 min after prompting. All questions are forced-choice, and are displayed as sequential screens on the smartphone. Participants will not be allowed to leaf through the present or previous diaries. The volume of the alarms is adjustable and there is the possibility to temporarily mute the alarm.

**Diary questionnaire**

The diary questionnaire consists of 32 items. In the morning and evening, five and eight additional questions are included, respectively. According to guidelines for diary questions, the items are short, simply worded, and try to mimic the participants’ internal dialogue, for example, ‘at the moment, I suffer from headache’.

**General health status and non-specific physical symptoms**

General health status will be assessed by using the first question from the RAND-36. Nine symptoms are selected that are most often reported by electro hypersensitive people according to studies in Switzerland and a survey among members of the Dutch Electrohypersensitivity Foundation. These symptoms comprise fatigue, distressed/nervous/tense feeling, concentration problems, tinnitus, dizziness or light-headedness, painful joints or muscles, skin problems, problems with vision, hearing or smell. In addition, one question asks for the symptom that the participant usually experiences and that is not in the prespecified list. For all symptoms the momentary experience is assessed (‘at this moment, …’). Response options range from ‘not at all’ to ‘very much’ on a five-point Likert response format with only the extremes labelled.

**Perceived exposure to (sources of) RF EMF, noise and air pollution**

Perceived exposure to RF EMF will be assessed both momentarily and for the interval between alarm cues, using the question ‘At the moment/since the last alarm cue, I am exposed to radio frequent electromagnetic fields’, with response options ranging from ‘not at all’ to ‘very much’ on a five-point scale with only the extremes labelled. Perceived exposure to specific sources of RF EMF is assessed by asking the participant to indicate which of the following sources mainly determined their exposure: mobile phone, DECT phone, WiFi, antennas for mobile telephony, radio or television masts, another source.

Momentary (but not for the interval between alarm cues) exposures to noise and air pollution are asked in a similar way.

**Environment**

In order to interpret the readings of the exposimeter, participants will be asked to indicate the kind of environment they were in during the interval between alarm cues. The environments included are at home inside, at home outside, elsewhere inside, elsewhere outside, on the road (on foot, by bike, car or public transport), in the city centre or a shopping area, in a residential or built-up area but not the centre, outside the built-up area (eg, in a rural area or in nature). All participants already completed a detailed time-activity questionnaire on their general behaviour in the EMPHASIS study.
Self-reported use of mobile phone and DECT phone

Phone use will be estimated separately for mobile phones and DECT phones based on the question of how many minutes a person called in the interval between alarm cues: 1–5, 5–10, 10–20, 20–30, 30–60 or longer than 60 min. In addition, participants will be asked to register on a form their use of mobile and DECT phones.

Mood

The Profile of Mood States (POMS) will be used to assess momentary (state) depression, vitality, anger and tension. Three items for each subscale will be used, as was carried out previously by Houtveen and van Doornen. The selected items were as follows: (1) depression: unhappy, sad, hopeless; (2) vitality: active, energetic, lively; (3) anger: angry, annoyed, moody and (4) tension: tense, nervous, anxious. Each item can be rated using a five-point Likert response format ranging from ‘not at all’ to ‘very much’ with only the extremes labelled.

Additional questions

After the morning alarm cue, the diary contains some questions about the duration and quality of sleep. Before the participants go to bed, questions are asked about use of medication to relieve their symptoms (mentioned in the diary), avoidance of sources of RF-EMF because of their symptoms, and whether the participants rested or took a nap during daytime.

Personal RF exposure assessment

Actual EMF exposure will be measured using EME-SPY 121 exposimeters (Satimo, Cortaboeuf, France) worn at the hip in a camera bag. As the maximum memory capacity of the exposimeter is 12,540 sampling intervals and the study will last for 120 h, the exposimeter will measure at an interval of 36 s. The exposimeters measure the RF electric field strength in 12 frequency bands used for communication and broadcasting (see online supplementary appendix 1). Before the exposimeters are employed in the study, they will be calibrated according to a previously described method, modified in a way that the calibrations take place in an anechoic chamber instead of a Gigahertz Transverse Electromagnetic cell (GTEM).

The participants will be instructed to wear the exposimeters all day except during wet activities (eg, showering) and sports (to avoid damage due to shocks). During sleep, the exposimeters have to be placed adjacent to the bed within 50 cm from the head. The electronic diary contains an event button that can be used to indicate when the exposimeters are worn or taken off.

GPS logger

Participants will wear a GPS device at their left shoulder. The GPS logger geo-locates the personal RF EMF measurements and the data can be used to visualise the participants’ location and measured EMF exposure on a Google Earth map. This visualisation can be used to interpret and check the quality of the EMF measurements.

Data analysis

Exposimeter data will be aggregated either over fixed time intervals or over intervals between random alarm cues, and for each interval the time weighted average, peak exposure, exposure above a certain threshold and rate of change can be calculated for all frequency bands separately and combined. The relationship of actual and perceived EMF exposure with non-specific physical symptoms will be analysed using multilevel regression analysis. The within-participant repeated measurements of actual and perceived EMF exposure and mood will be the first-level variables, which will be modelled as fixed effects; the second level will be the individuals and will be modelled as random effects. As time (hour of the day) is associated with symptom occurrence and severity (eg, for fatigue), this will be included in the model using a sinus-cosinus 24 h function to account for the diurnal pattern. Unique contributions of the various explanatory variables will be estimated in multivariate models, in which an intercept, 24 h time function, actual EMF exposure, perceived EMF exposure and mood will be included. Time-shift models with different lag times will be used to gain insight into the time delay between exposure and mood and the occurrence of symptoms. The time window of 5 days should be sufficient to capture the potential participants’ response to EMF exposure, based on reported latencies between exposure and symptoms.

A multilevel power analysis was performed to calculate the strength of the association between perceived EMF exposure and non-specific physical symptoms that could be detected with the repeated measurements multilevel analysis in a sample of 60 volunteers with each 40 (5 days with 8 measurements) observations. A significance level of 5% and a power of 80% were used. This method based on simulations has been described previously. No such calculation was made for actual EMF exposure because most evidence points towards the absence of an association with non-specific physical symptoms.

Input parameters for the power analysis came from a pilot study of four (not electrohypersensitive) master students (2 men and 2 women) who completed the diary questions for 2 weeks (unpublished data, National Institute for Public Health and the Environment 2012). Figure 1 illustrates how the power of the statistical analysis differs according to the magnitude of the regression coefficient. It can be seen that at a power of 80%, the detectable regression coefficient is slightly over 1.5, which corresponds to an increase of 1.5 on the sum of 10 symptoms (range 0–40) at an increase of 1 in perceived exposure (range 0–4)
been shown that especially in people with high levels of
non-specific physical symptoms, less symptoms are
reported when using momentary assessment methods
compared with retrospective methods. Since persons
with IEI-EMF typically report more symptoms than the
general population, a symptom diary is more suitable
to obtain a valid estimate than asking to report the usual
or average number of symptoms retrospectively.

Regarding the EMA measurement of exposure, exposi-
metrys are the method of choice to measure personal
exposure compared with spot measurements, self-
estimated exposure and exposure prediction models.
Nevertheless, measurements of exposimeters also have
their own limitations and are not always free of bias. To
obtain valid measurements, it is important that the par-
ticipants receive clear and standardised instructions about
how to wear the metres. In this way, measurements by
different participants will be harmonised and more com-
parable. Exposure from mobile phone use by the partici-
pants themselves will not be measured properly by the
exposimeter because the exposimeters are designed to
measure the far EMF field, that is, exposure from EMF
sources further away. In the near field, the exposimeters
are unable to correctly measure the exposure, resulting
in overestimations or ‘clipping’ in which the maximum
measurement value of 10 V/m is registered. Therefore,
mobile phone use during each 2.5 h interval between
alarm cues is asked for in the diaries (although we are
aware of the recall bias in the estimates of self-reported
phone use) and participants are requested to register
their use of mobile and DECT phones.

Another strength of using individual-level exposure
and outcome data is that confounding by factors that
remain stable over time is reduced because within-
participant variation in exposure and manifestation of
health symptoms are of primary interest. Thus, the ana-
lyses do not require adjustment for personal characteris-
tics such as demographic factors and psychological traits.
However, to detect possible interaction effects, for
example, different associations in men and women, per-
sonal characteristics have to be taken into account. The
multilevel regression analysis allows for such an investiga-
tion of cross-level interactions. As the study population
may be a mix of ‘truly’ electrosensitive individuals and
individuals in whom psychological mechanisms account
for their symptoms, we will explore the applicability of
statistical methods to study associations that differ
between individuals (vector autoregressive models).

Time-varying factors associated with EMF exposure
that potentially affect symptom occurrence also need to
be accounted for. Avoidance of EMF exposure after
symptoms begin to develop is such a factor (which is
asked for in the diary). Time-varying factors associated
with symptoms but not with EMF exposure are no con-
founders, but can obscure relationships between expos-
ure and symptoms. This is the reason a question about
the use of medication to relieve symptoms and rest/sleep
during the day is included in the diary questionnaire.

ETHICS AND DISSEMINATION

A formal written inquiry, including a detailed descrip-
tion of the study protocol, was made at the Medical
Ethics Committee of the University Medical Centre
Utrecht to verify whether the study protocol should be
tested within the framework of the Dutch Medical
Research Involving Human Subjects Act. A formal
advice was received in which the Committee indicated
that the study was exempt from having to pass the full
ethics testing procedure. Handling of personal data will
comply with the Personal Data Protection Act (in Dutch:
Wet bescherming persoonsgegevens (Wbp)). After
removal of the identifying information, data will be
stored on a part of the institute’s server with limited
access by specified employees.

Results of the study will be offered for publication in
the international peer-reviewed literature and presented
at (international) conferences. Further, results will be
disseminated at a national level at meetings organised by
The Netherlands Organisation for Health Research and
Development (ZonMw). At these meetings representa-
tives of IEI-EMF patient groups will also be reached.

DISCUSSION

The EMA design of this study is innovative as it com-
bines actual exposure measurements with momentary-
measured health symptoms. The design aims to circum-
vent important limitations of previous studies into
IEI-EMF, which can be summarised as biased recall of
health outcomes, poor characterisation of individual
exposure to EMF and experimental exposure conditions
that substantially differ from the real-life situation (ie,
low ecological validity). Another potential strength of
the study is the simultaneous assessment of actual and
perceived EMF exposure. In the multilevel regression
analysis, independent contributions of these two types of
exposure can be estimated, an approach similar to the
EMPHASIS study. With respect to recall bias, it has
been shown that especially in people with high levels of

Figure 1  Power of the repeated measurements multilevel
regression analysis as a function of the magnitude of the
regression coefficient, n=60 persons with 40 repeated
measurements.
There are also limitations to the described study. Although the external (ecological) validity of the design is high, compared with double-blind trials such as provocation studies in the laboratory, the internal validity is lower. Further, the study involves relatively high costs in terms of the necessary equipment and work. The exposimeters are expensive and vulnerable to physical damage, and the electronic diaries are also costly compared with traditional paper and pencil diaries. Also, the programming of the diary software is very time consuming. As a result, only a limited number of participants can be included in the study. The burden for participants is quite high, since they have to carry the exposimeters and GPS loggers with them and are interrupted by the diary alarm cues several times a day. It is expected that only highly motivated persons will participate, but because the study is not intended to examine a representative sample of the population this is not considered a problem. Compliance to the study protocol may be difficult for some participants because they may find wearing the exposimeter awkward. As a result of participation in the study, the participants’ attention to EMF exposure and physical symptoms may increase (reactivity). It can in principle not be excluded that the data are manipulated by placing the metres adjacent to a (assumed) source of EMF before an alarm cue is expected. To minimise the chance of such anticipation effects, the alarms are programmed at random intervals. Further, unusual exposure patterns can be checked for, although high and prolonged exposure peaks can be real and do not have to result from anticipation effects.

In the statistical analysis, false-positive associations due to multiple testing may arise. For EMF exposure, there is information about 12 frequency bands, which can be analysed separately and combined. Possible exposure metrics include time-weighted average, peak exposure, rate of change and exposure above a certain threshold. Since no biological mechanism is known that explains how EMF can affect health, in theory each combination of almost 50 combinations of frequency and exposure metrics can be relevant. To take into account the possibility of false-positive associations, the expected proportion of falsely rejected hypotheses will be controlled using a sequential Bonferroni-type procedure described by Benjamini and Hochberg. If associations with symptoms are detected, it is necessary to replicate the results to exclude chance. Therefore, by excluding persons based on psychological characteristics the study population of interest may be missed. Somatic (chronic) diseases will not be excluded either, as a considerable part of persons having IEI-EMF suffer from chronic diseases, and at this stage it is not possible to decide on theoretical grounds which medical conditions should be included or excluded.

A control group is not deemed necessary, since in the longitudinal study design each participant acts as their own control. Moreover, if there are associations between RF EMF exposure and symptoms, this will be most likely in persons who report to be sensitive to RF EMF, and statistical power will be enhanced by focusing on this group. A limiting factor of this approach is that we cannot verify if persons with IEI-EMF are more sensitive to EMF than controls.

In summary, this is the first time that actual and perceived exposure and possibly explanatory variables are combined to such extent in an IEI-EMF sample. The main strengths of the study described in this article are an accurate assessment of EMF exposure and non-specific physical symptoms using an EMA methodology, and elimination of confounding by personal characteristics as a result of the within-person design. The methodological issues aforementioned will be accounted for and limited as much as possible in the conduct of the study as well as the analysis and interpretation of its outcomes.

**Contributors** RPB conceived the study, participated in its design and coordination, and drafted the manuscript. JFBB, JHH, RTvS, CAMS and CB participated in the study design and helped to revise the manuscript critically for important intellectual content. EL conceived the study, participated in the study design and helped to revise the manuscript critically for important intellectual content. MA programmed the electronic diary software and helped to revise the manuscript critically for important intellectual content. IVK participated in the study design, permitted access to the EMPHASIS dataset and helped to revise the manuscript critically for important intellectual content. All authors read and approved the final manuscript.

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