

SYSTEMATIC REVIEW

Mobile phones affect multiple sperm quality traits: a meta-analysis [version 1; referees: 2 approved, 1 approved with reservations]

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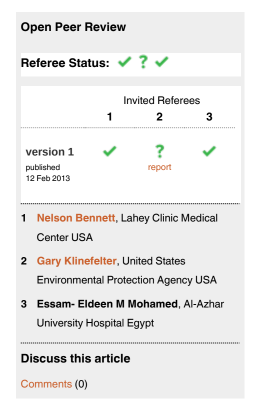


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Abstract

As mobile phone usage is growing rapidly, there is a need for a comprehensive analysis of the literature to inform scientific debates about the adverse effects of mobile phone radiation on sperm quality traits. Therefore, we conducted a meta-analysis of the eligible published research studies on human males of reproductive age. Eleven studies were eligible for this analysis. Based on the meta-analysis, mobile phone use was significantly associated with deterioration in semen quality (Hedges's g = -0.547; 95% CI: -0.713, -0.382; p < 0.001). The traits particularly affected adversely were sperm concentration, sperm morphology, sperm motility, proportion of non-progressive motile sperm (%), proportion of slow progressive motile sperm (%), and sperm viability. Direct exposure of spermatozoa to mobile phone radiation with in vitro study designs also significantly deteriorated the sperm quality (Hedges's g = -2.233; 95% CI: -2.758, -1.708; p < 0.001), by reducing straight line velocity, fast progressive motility, Hypo-osmotic swelling (HOS) test score, major axis (µm), minor axis (μ m), total sperm motility, perimeter (μ m), area (μ m²), average path velocity, curvilinear velocity, motile spermatozoa, and acrosome reacted spermatozoa (%). The strength of evidence for the different outcomes varied from very low to very high. The analysis shows that mobile phone use is possibly associated with a number of deleterious effects on the spermatozoa.



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Introduction

Almost 10% of men of reproductive age are estimated to be subfertile¹. Owing to its complexity, even after identification of a plethora of underlying factors, etiology in almost half of the infertile subjects tested at fertility clinics remains obscure². Hence, the list of the causes of male infertility is growing by the day with recent advances in fertility research³. Though advances in assisted reproduction technologies (ARTs), especially in the form of *in vitro* fertilization (IVF) and intracytoplasmic sperm injection (ICSI), have helped subfertile couples conceive offspring, it is feared that ARTs only bypasses the problem of subfertility and contributes towards hiding the underlying causes which have at times led to serious health problems in offspring^{4,5}. Hence, identification of unknown aetiologies would help in prescription of specific preventive measures that will ultimately decrease the incidence of male infertility.

Most nations, especially developing countries, are witnessing an increase in the use of various radiation-emitting domestic-purpose devices that could cause mild to serious health problems based on the duration and intensity of usage⁶, and reduced fertility is now recognised as one such problem7. Wireless mobile phones are one of the most accepted devices with a tremendous increase in usage across the world in recent times⁸. Research into the impact of ionizing radiation on the development of various types of health disorders, especially cancers, has been well established⁹. Similarly, several studies have found an increase in the risk of developing some types of tumors after long-term exposure to non-ionizing radiation from mobile phones¹⁰. Research into the effects of mobile phone radiation on male fertility, though growing, is limited and inconclusive^{11,12}. Recently, several case-control studies have reported results from a general population setting alongside a few studies from subfertile populations^{7,13–20}. Like ionizing radiation, non-ionizing radiation is also expected to affect spermatozoa, though in subtle ways²¹. The aim of this meta-analysis was, therefore, to investigate the impact of mobile phone radia in vitro as well as in vivo settings in men of reproductive age from both general and subfertile populations.

Material and methods

A systematic search of an electronic database was conducted to retrieve published studies on the impact of mobile phone radiation on semen parameters in adult men. The results have been reported according to the standards of the guidelines for meta-analysis of observational studies in epidemiology²². All English language research studies published up until January 2012 in scientific journals indexed in the searched databases were included for analysis.

Inclusion/exclusion criteria and outcomes of interest: The studies on human males of reproductive age reporting the effect of mobile phone radiation on any or all measures of semen volume, total sperm count, sperm concentration, sperm motility or sperm morphology were included. All the studies that did not satisfy the inclusion criteria were excluded.

Search strategy, data extraction and meta-analysis: Google Scholar and NLM's PubMed database were searched for articles by using different combinations of 4 mobile phone related keywords ['mobile

phone', 'cellular phone', 'radiofrequency electromagnetic waves (RF-EMW)', 'radiation'] with 5 sperm quality related keywords ('spermatozoa', 'semen', 'sperm concentration', 'sperm motility', 'sperm morphology') Data from 11 eligible studies were extracted and separated into *in vitro* and *in vivo* categories.

Effect sizes were expressed as Hedges's g²³, separately for *in vivo* & *in vitro* studies using individual semen parameters as units of analysis (Supplementary Table 1). A random model was used to test and quantify effect size using 'Comprehensive Meta-Analysis (v.2)' trial version²⁴. A random effect model was preferred over a fixed effect model in order to account for differences in both effect size and sampling error²⁵.

Results

In vivo effects of mobile phone radiation

Our analysis shows that overall, mobile phone users had significant deterioration in semen quality (Hedges's g = -0.547; 95% CI: -0.713, -0.382; p < 0.001). There was significant heterogeneity among effect sizes (Q = 475.985, p < 0.001), which suggest that some of the semen parameters may not be affected by mobile phone exposure. Hence, combined effect-size for each of the semen parameters were calculated separately (Table 1), and it was found that sperm concentration, sperm morphology, sperm motility, proportion of non-progressive motile sperm (%), proportion of slow progressive motile sperm (%), and sperm viability were deteriorated in individuals exposed to mobile phone radiation. By contrast, semen volume, liquefaction time, semen pH, proportion of rapid progressive motile sperm (%), and semen viscosity were not affected by mobile phone usage.

Publication bias could potentially change the results of metaanalysis but analysis of funnel plot of precision by Hedges's g using Dual and Tweedie's trim-and-fill test²⁶ did not change the overall effect size, suggesting little bias. Moreover, Rosenthal's fail-safe N test²⁷ revealed that 3964 missing studies with a mean Hedges's g of 0 are required for the combined 2-tailed p-value to exceed 0.050. In other words, there need to be 99.1 missing studies for every observed study for the effect to be nullified.

In vitro effects of mobile phone radiation

Experimental exposure of spermatozoa isolated from healthy men of reproductive age to mobile phone radiation significantly affected sperm quality (Hedges's g = -2.233; 95% CI: -2.758, -1.708; p < 0.001). There was significant heterogeneity among effect sizes (Q = 639.294, p < 0.001), suggesting that similar to *in vivo* exposure, in vitro exposure may also not affect all the parameters of spermatozoa. Hence, combined effect-size for spermatozoa parameters were calculated separately (Table 1), and it was found that exposure to mobile phones significantly reduced straight line velocity, fast progressive motility, Hypo-osmotic swelling (HOS) test score, major axis (µm), minor axis (µm), total sperm motility, perimeter (µm), area (µm²), average path velocity, curvilinear velocity, motile spermatozoa, and acrosome reacted spermatozoa (%). By contrast, DNA fragmentation levels, non-progressive motility, total antioxidant capacity (TAC), progressive motility, reactive oxygen species (ROS) generation, slow progressive motility, sperm concentration, and sperm zona binding was not affected by mobile phone radiation.

Table 1. Effect sizes of mobile phone radiation on sperm quality traits.

	Sample size	Hedges's g	p-value
In vivo stu	udies		
Semen volume	591	0.09774	0.29458
Sperm concentration	874	-0.66388	0.01858
Sperm morphology	746	-1.28325	0.00000
Sperm motility	1079	-0.81584	0.00102
Proportion of non-progressive motile sperm (%)	283	-0.16136	0.03396
Proportion of rapid progressive motile sperm (%)	283	-0.25708	0.09969
Proportion of slow progressive motile sperm (%)	283	-0.39031	0.00765
Liquefaction time (min)	321	-0.11449	0.28277
pH	321	-0.36681	0.05592
Sperm viability (%)	321	-1.13150	0.00220
Semen viscosity	321	-0.00924	0.93083
In vitro sto	udies		
Acrosome reaction (%)	24	-1.69939	0.00000
Sperm area (µm²)	24	-6.79952	0.00004
Average path velocity	20	-8.16777	0.00000
Curvilinear velocity	20	-10.37987	0.00000
DNA fragmentation	32	0.10182	0.68034
Fast progressive motility	49	-0.50794	0.01195
Hypo-osmotic swelling (HOS)	20	1.721867	0.000002
Major axis (µm)	24	-3.62708	0.01918
Minor axis (µm)	24	-7.4825	0.0361
Sperm motility	105	-2.82739	0.00118
Non motile spermatozoa	49	-0.61615	0.03275
Non progressive motility	49	0.04371	0.82612
Perimeter (µm)	24	-5.53132	0.01897
Progressive motility	12	-0.04606	0.90700
Reactive oxygen species (ROS)	36	-11.37087	0.33592
Slow progressive motility	49	-0.14543	0.67535
Sperm concentration	59	-0.02309	0.89887
Sperm zona binding	10	-0.68402	0.12153
Straight line velocity	20	-6.37614	0.00000
Total antioxidant capacity (TAC)	32	-0.25102	0.31138
Viability (%)	56	-2.75116	0.02543

A Funnel plot of precision by Hedges's g using Dual and Tweedie's trim-and-fill test did not change the overall effect size, suggesting little publication bias. Rosenthal's fail-safe N test revealed that 3813 missing studies with a mean Hedges's g of 0 are required for the combined 2-tailed p-value to exceed 0.050. In other words, there need to be 100.3 missing studies for every observed study for the effect to be nullified.

Discussion

This study was aimed to analyse the data assessing the risk of mobile phone radiation on male fertility. Our results suggest that mobile phone radiation has a tendency to significantly affect sperm quality. Based on the design of the analysed records, we divided studies into *in vivo* studies and *in vitro* studies. The effect size was significant in both the categories, suggesting that mobile phone

radiation could severely compromise male fertility. This conclusion is robust, as a fail-safe test suggested that the results are not likely to be mediated by publication bias.

The number of worldwide mobile subscriptions grew from less than 1 billion in 2000 to over 6 billion in 2012⁸, with more than half of these subscribers estimated to be children and young adults. Hence, it is very likely that in the coming decades, we could witness an increase in the incidence of male infertility due to mobile phone radiation exposure, similar to growing concerns over other hazards. Although the mechanism of cell phone radiation-mediated health defects is still obscure, it is proposed that their ability to produce heat, disrupt cell membranes, affect endothelial function, alter the blood-brain barrier, and modulate neuronal excitability have the potential to affect multiple physiological functions simultaneously^{28–30}.

To our knowledge, this is the first meta-analysis of the effects of mobile phone radiations on various sperm quality parameters. Cellular phones have become integral part of everyday life, and newer versions of these are developed very rapidly these days. Hence, it is necessary to educate the users about the hazards of cell phones as well as test the newer versions like smartphones for health hazards.

Competing interests

No relevant competing interests were disclosed.

Grant information

The author(s) declared that no grants were involved in supporting this work.

Supplementary table

Supplementary Table 1. Effect sizes of sperm quality traits from the studies included in the analysis.

Reference	Subgroup	Outcome	Effect size (Hedges's g)	p-value			
In vivo studies							
[7]		Proportion of non-progressive motile sperm (%)	-0.11444	0.19545			
		Proportion of rapid progressive motile sperm (%)	-0.39434	0.00001			
	1	Proportion of slow progressive motile sperm (%)	-0.51671	0.00000			
		Sperm concentration	0.01922	0.82778			
		Sperm motility	-0.14692	0.09667			
	2	Proportion of non-progressive motile sperm (%)	-0.28478	0.04855			
		Proportion of rapid progressive motile sperm (%)	-0.07945	0.58037			
		Proportion of slow progressive motile sperm (%)	-0.22090	0.12525			
		Sperm concentration	-0.12467	0.38594			
		Sperm motility	0.00940	0.94784			
5.403	1	Sperm morphology	-0.74105	0.00000			
[13]		Sperm motility	-0.57347	0.0000			
		Liquefaction time (min)	-0.01209	0.94773			
		Hq	0.00000	1.00000			
		Semen volume	0.18269	0.32253			
		Sperm concentration	-0.42958	0.02095			
	1	Sperm morphology	-0.72462	0.00013			
		Sperm motility	-0.40596	0.02896			
		Viability	-0.43282	0.02002			
		Viscosity	0.01942	0.91612			
	2	Liquefaction time (min)	-0.23709	0.20389			
		pH	-0.46407	0.01363			
		Semen volume	-0.02014	0.91380			
E4.43		Sperm concentration	-0.56141	0.00298			
[14]		Sperm morphology	-1.70950	0.0000			
		Sperm motility	-1.32047	0.0000			
		Viability	-1.34677	0.00000			
		Viscosity	-0.09456	0.61148			
		Liquefaction time (min)	-0.09749	0.59412			
	3	Hq	-0.63951	0.00060			
		Semen volume	0.28711	0.11786			
		Sperm concentration	-0.87694	0.00000			
		Sperm morphology	-1.95983	0.0000			
		Sperm motility	-1.58904	0.0000			
		Viability	-1.62719	0.00000			
		Viscosity	0.04490	0.80606			
	1	Semen volume	-0.07567	0.69348			
5453		Sperm concentration	-2.09426	0.00000			
[15]		Sperm morphology	-1.35171	0.00000			
		Sperm motility	-1.80265	0.00000			
		Overall effect	-0.54948	0.00000			
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Reference	Subgroup	Outcome	Effect size (Hedges's g)	p-value		
In vivo studies						
[16]		Fast progressive motility	-0.48612	0.07419		
		Motility	-0.73467	0.00808		
	1	Non motile	-0.89668	0.00146		
	I	Non progressive motility	0.14043	0.60105		
		Slow progressive motility	-0.48268	0.07620		
		Sperm concentration	-0.05135	0.84822		
	1	Dna fragmentation	0.10182	0.68034		
		Motility	-0.19307	0.43544		
[00]		ROS	-0.29465	0.23542		
[20]		Sperm concentration	0.00085	0.99725		
		TAC	-0.25102	0.31138		
		Viability (%)	-0.46743	0.06193		
[17]	1	Progressive motility	-0.04606	0.90700		
	1	Motility	-16.10595	0.00008		
[31]		ROS	-23.97770	0.00007		
		Viability (%)	-11.52174	0.00009		
		Acrosome (%)	-1.58348	0.00051		
		Area (µm²)	-8.61098	0.0000		
	1	Major axis (µm)	-5.25493	0.00000		
		Minor axis (µm)	-11.21546	0.00000		
		Perimeter (µm)	-8.00952	0.0000		
[32]		Sperm zona binding	-0.68402	0.12153		
	2	Acrosome (%)	-1.82487	0.00012		
		Area (µm²)	-5.27741	0.0000		
		Major axis (µm)	-2.15357	0.00002		
		Minor axis (µm)	-4.06799	0.00000		
		Perimeter (µm)	-3.28849	0.0000		
[18]	1	Fast progressive motility	-0.53471	0.07618		
		Motility	-0.64188	0.03467		
		Non motile	-0.31928	0.28406		
		Non progressive motility	-0.07395	0.80286		
		Slow progressive motility	0.21209	0.47510		
	1	Average path velocity	-8.16777	0.0000		
[19]		Curvilinear velocity	-10.37987	0.00000		
		HOS	1.72187	0.00000		
		Motility	-9.78102	0.0000		
		Straight line velocity	-6.37614	0.00000		
		Viability (%)	-2.53934	0.00000		
		Overall effect size	-2.23292	0.00000		
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Open Peer Review

Current Referee Status:







Version 1

Referee Report 25 March 2013

doi:10.5256/f1000research.862.r861



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Department of Dermatology and Andrology, Al-Azhar University Hospital, Assiut, Egypt

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.

Referee Report 14 March 2013

doi:10.5256/f1000research.862.r808



Gary Klinefelter

Gamete and Early Embryo Biology Branch, Reproductive Toxicology Division, United States Environmental Protection Agency, Research Triangle Park, NC, USA

On the surface, the results seem quite striking with virtually any sperm endpoint one can imagine being significantly altered in the collective analysis of mobile phone studies compiled. However, upon looking at the data in the supplementary table, it is obvious that the relatively few studies compiled varied widely both in respect to endpoints measured and the sample size. As shown in Table 1, motility is the endpoint representing the greatest combined sample size for both *in vivo* and *in vitro* studies. Motility was measured in 4 out of 4 *in vivo* studies and 5 out of 7 *in vitro* studies. So motility 'might' be an endpoint that is repeatedly altered by cell phone exposure. The reason for 'might' is the lack of any reported exposure data in this study.

In summary, the small sample size and lack of exposure data significantly weaken the conclusions of this study.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Competing Interests: No competing interests were disclosed.

Author Response 02 Apr 2013



Madhukar Dama, IWVR, KVAFSU, India

We have studied the reviewer comments and would like to justify our results. Our analysis is showing that mobile phone radiations could affect many sperm parameters. This could be due to interdependence of sperm parameters (Acta Eur Fertil. 1982;13(2):49-54). We also agree with the point that the number of studies is few and total sample size in *in vitro* studies is smaller. However, it must be noted that the sample size is weighted during meta-analysis, which nullifies the problems posed by smaller sample size studies. Apart from motility, other parameters like morphology, concentration, and viability are also significantly affected by *in vivo* exposure. Hence we have provided all the effect sizes individually along with p values and sample size. We hope that our points justify the reviewer comments.

Competing Interests: None

Referee Report 19 February 2013

doi:10.5256/f1000research.862.r780



Nelson Bennett

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I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.