



2 Elevated silver, barium and strontium in 3 antlers, vegetation and soils sourced from 4 CWD cluster areas: Do Ag/Ba/Sr 5 piezoelectric crystals represent the 6 transmissible pathogenic agent in TSEs?

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Summary High levels of Silver (Ag), Barium (Ba) and Strontium (Sr) and low levels of copper (Cu) have been measured in the antlers, soils and pastures of the deer that are thriving in the chronic wasting disease (CWD) cluster zones in North America in relation to the areas where CWD and other transmissible spongiform encephalopathies (TSEs) have not been reported. The elevations of Ag, Ba and Sr were thought to originate from both natural geochemical and artificial pollutant sources – stemming from the common practise of aerial spraying with ‘cloud seeding’ Ag or Ba crystal nuclei for rain making in these drought prone areas of North America, the atmospheric spraying with Ba based aerosols for enhancing/refracting radar and radio signal communications as well as the spreading of waste Ba drilling mud from the local oil/gas well industry across pastureland. These metals have subsequently bioconcentrated up the foodchain and into the mammals who are dependent upon the local Cu deficient ecosystems. A dual eco-prerequisite theory is proposed on the aetiology of TSEs which is based upon an Ag, Ba, Sr or Mn replacement binding at the vacant Cu/Zn domains on the cellular prion protein (PrP)/sulphated proteoglycan molecules which impairs the capacities of the brain to protect itself against incoming shockbursts of sound and light energy. Ag/Ba/Sr chelation of free sulphur within the biosystem inhibits the viable synthesis of the sulphur dependent proteoglycans, which results in the overall collapse of the Cu mediated conduction of electric signals along the PrP-proteoglycan signalling pathways; ultimately disrupting GABA type inhibitory currents at the synapses/end plates of the auditory/circadian regulated circuitry, as well as disrupting proteoglycan co-regulation of the growth factor signalling systems which maintain the structural integrity of the nervous system. The resulting Ag, Ba, Sr or Mn based compounds seed piezoelectric crystals which incorporate PrP and ferritin into their structure. These ferrimagnetically ordered crystals multireplicate and choke up the PrP-proteoglycan conduits of electrical conduction throughout the CNS. The second stage of pathogenesis comes into play when the pressure energy from incoming shock bursts of low frequency acoustic waves from low fly jets, explosions, earthquakes, etc. (a key eco-characteristic of TSE cluster environments) are absorbed by the rogue ‘piezoelectric’ crystals, which duly convert the mechanical pressure energy into an electrical energy which accumulates in the crystal-PrP-ferritin aggregates (the fibrils) until a point of ‘saturation polarization’ is reached. Magnetic fields are generated on the crystal surface, which initiate chain reactions of deleterious free radical mediated spongiform neurodegeneration in surrounding tissues. Since Ag, Ba, Sr or Mn based piezoelectric crystals are heat

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resistant and carry a magnetic field inducing pathogenic capacity, it is proposed that these ferroelectric crystal pollutants represent the transmissible, pathogenic agents that initiate TSE.

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39 Introduction

40 Exceptionally high levels of Ag, Sr and Ba were
41 measured in deer antlers, vegetation, soils sourced
42 from chronic wasting disease (CWD) cluster areas in
43 Colorado, Wisconsin, Saskatchewan, whilst levels
44 were 3-fold, 2.5-fold and 3-fold less (for Ag, Sr, Ba,
45 respectively) in CWD-free areas of Alberta/UK. Ag
46 was virtually undetectable in antlers sampled from
47 CWD-free deer herds in the UK.

48 These observations were recorded as part of an
49 extensive comparative analytical study of the
50 levels of 46 metals in the soils, water and vege-
51 tation of CWD cluster and CWD-free regions con-
52 ducted across North America. This work
53 represented the North American perspective of a
54 three year globally orientated project designed to
55 establish whether any abnormal mineral profile or
56 abnormal magnetic/radioactive/oxidative capac-
57 ity is a common characteristic of transmissible
58 spongiform encephalopathie (TSE) cluster ecosys-
59 tems around the world, and, if so, whether that
60 abnormality plays a primary role in the patho-
61 genesis of TSEs.

62 Since the primary origins of TSEs are unknown,
63 this study was designed to challenge the theory
64 based upon previously amassed data [1–3] that
65 high levels of specific metals, such as manganese
66 (Mn) or Ag, in combination with low levels of Cu in
67 the environment may bring about a rogue metal
68 replacement at vacant Cu ligands on the cellular
69 prion protein (PrPc) – a Cu binding protein [4]
70 whose misfolded isoforms hallmark the TSE dis-
71 eased brain [5].

72 The Hypothesis: Pt 1, the healthy func- 73 tion of Cu–PrP

74 Does a network of Cu–PrP–Cu-proteoglycan 75 conduits conduct a relay of electric signals 76 that regulate the auditory/circadian 77 associated circuitry of the biosystem?

78 Since PrPc ‘knock out’ mice develop symptoms
79 (sleep disorders, abnormal EEG, etc.) that indicate

an underlying disturbance in the regulation of the 80
nocturnal–diurnal rhythm [6,7] and GABA inhibi- 81
tory currents at synapses [4,5], it has been sug- 82
gested that PrPc performs some metabolic role in 83
mediating the circadian diurnal–nocturnal rhythm 84
and other external stimuli. In this respect, it has 85
been proposed that the paramagnetic Cu [8] com- 86
ponent of the PrP molecule [4] performs a role as a 87
conductor of electrical signals [2,3] that mediate 88
the neuronal response to external diurnal–noctur- 89
nal, auditory, tactile stimuli, etc.; whereby the 90
electrical signals that are transduced from the in- 91
coming energy of these external stimuli, are re- 92
layed to the synapses for regulating the GABA 93
inhibitory amino acids/excitatory amino acids that 94
ultimately modulate the neuronal response. It is 95
interesting that PrPc and copper are highly con- 96
centrated at the synapse/end plates of select 97
neuronal tracts in the CNS [4], whilst PrP has been 98
shown to influence GABA-type inhibitory currents 99
[4]. 100

Ultraviolet, acoustic and other sources of in- 101
coming energy enter the organism and are trans- 102
duced at the melanin granules (located in the 103
retina, skin [9] and cochlear cells [10]) into electric 104
signals which are conducted via Cu atoms onto 105
PrP’s octapeptide repeat metallo binding domains, 106
whereupon that electrical energy is relayed from 107
prion protein to prion protein – ‘domino-style’ – 108
to synapses/end plates along the auditory, vestib- 109
ular and circadian associated circuitry for 110
modulating the GABA type inhibitory currents, 111
which, in turn, modulates the overall neuronal re- 112
sponse to the original external stimulus. In this 113
respect, these electrical signals perform a key 114
communicative role in connecting the external di- 115
urnal/nocturnal, auditory, tactile stimuli with a 116
broad diversity of circadian/auditory effector tis- 117
sues/organs throughout the biosystem. 118

The Paramagnetic Cu co-partners of PrP could 119
be viewed as conducting the electrical signals into 120
the synapses/endplates in order to modulate the 121
wide array of physiological processes under audi- 122
tory, tactile, diurnal–nocturnal circadian regula- 123
tion – e.g., sleep/wake rhythms, sexual cycles, 124
mood/behaviour, heart beat, immune response, 125
gastrointestinal rhythms, growth and repair of cells 126

127 (particularly during embryonic/early develop-
128 ment), including the growth of tumour cells
129 [11–14].

130 Interestingly, PrPc is intensively localised/ex-
131 pressed in these neural/extra neural tissues that
132 are directly associated with the circadian/audi-
133 tory circuits and/or their target tissues – e.g., in
134 the cochlear [15], retina, pineal, hypothalamus,
135 visual cortex, pituitary, medulla, glial cells, sym-
136 pathetic neurones, spleen, lymphatic, tonsils,
137 appendix, myocardial cells, nerve growth factor
138 (maintaining growth and repair) mediated tissues
139 [5,16–18].

140 It is possible that the Cu component of the PrP
141 delivers its electrical ‘cargo’ to a copper centred
142 sulphated proteoglycan molecule [19,20] and that
143 these two molecules may play some broad ranging
144 ‘two way’ co-operative role in the conduction of
145 electrical signals around the biosystem; where the
146 Cu components on PrP deliver their ‘spark’ to the
147 sulfated proteoglycans (heparin) – growth factor
148 co-receptors [21]; which, in turn, regulate the all
149 important signalling system that maintains/modu-
150 lates the growth and structural integrity of the
151 nervous system [21]. It is interesting that proteo-
152 glycans are known to bind with copper [19] as well
153 as PrPc [5,22,23] in the healthy mammalian bio-
154 system, and that the successful binding of PrP to
155 proteoglycans is disrupted in the TSE diseased or-
156 ganism [23].

157 Further support for these ideas on PrPc’s healthy
158 role can be gleaned from the clinical and patho-
159 logical profiles that emerge from the disruption/
160 loss of PrP function in TSEs. For instance, the initial
161 lesions in TSEs tend to erupt in the retina [24,25],
162 the skin and the cochlear [15] – e.g., cell lines
163 where the melanin ‘transducer stations’ are in-
164 tensively expressed [10,11] – involving tissues that
165 are in the front line of receiving incoming light,
166 tactile and sound stimuli. Many of the more distal
167 regions of the auditory, optic and vestibular tracts
168 are also lesioned in TSEs [26]. Furthermore, one of
169 the key early stage clinical features in BSE suffering
170 cattle is involves a pronounced ‘hypersensitivity’
171 to sound, touch and light. UK vets actually ex-
172 ploited the symptom of hyperacusis in BSE, by ap-
173 plying a simple ‘handclap’ startle response test as
174 the best means of diagnosing clinical BSE.

175 Interestingly, genetically engineered mutations
176 in the melanocyte cells (the melanin producing
177 cells [11]) of experimental animals have been
178 shown to induce spongiform encephalopathy [27];
179 suggesting that a disruption at any point along
180 these putative conduits of electrical signalling
181 could be associated with the initiation of TSE
182 pathogenesis.

**The Hypothesis: Pt 2, metal nucleated
crystals and the unhealthy dysfunction
of PrP** 183
184
185

**Ag, Ba, Sr or Mn nucleated crystal-PrP-
ferritin complexes disrupt these conduits of
electrical signalling throughout the CNS.
A primary prerequisite for the pathogenesis
of TSEs? (see Fig. 1)** 186
187
188
189
190

In this respect, the pathogenesis of TSE could be 191
initiated by a disruption at any point along these 192
putative PrP-proteoglycan ‘conduits’ of electrical 193
conduction; ultimately resulting in the overall 194
collapse in both the GABA mediated modulation of 195
neuronal response at the synapses [4] and the 196
proteoglycan regulated anti-oxidant/growth factor 197
signalling systems [21,28]. 198

Environmental analytical observations in TSE 199
cluster zones suggest that a foreign metal re- 200
placement binding on the native Cu/Zn domains 201
on PrPc [4] and/or the proteoglycans [19,20] 202
could be implicated here. This would involve a 203
substitution by a rogue ferrimagnetic or diamag- 204
netic metal species that disrupts the normal 205
paramagnetic conductive capacity of PrP’s Cu co- 206
partner. It is also possible that a radioactive metal 207
species could be involved as the rogue metal re- 208
placement here. 209

So once the vacant Cu domains on the prion 210
protein/proteoglycans have been substituted by a 211
rogue reactive metal species, it is easy to envision 212
how the brain could be subjected to a steady, self 213
perpetuating state of deleterious free radical ‘melt 214
down’. For these foreign substitute metals will fail 215
to act in the overall best interests of the organism 216
and conduct electric signals in a balanced way. 217

**Rogue metals chelate sulphur and inactivate S-
proteoglycans?** 218
219

Chronic exposures to the highly reactive Ba, Sr salts 220
are known to invoke various other metabolic dis- 221
turbances [43] that are evident in the pathogenesis 222
of TSEs – such as the hyperactivation of calcium 223
and potassium channels [5]. The fact that Ba, Sr, 224
Ag and Mn will readily conjugate with sulphur in the 225
biosystem [43] could be very pertinent to TSEs, 226
since this would deprive the endogenous sulph- 227
ated(S)-proteoglycan molecules of their crucial 228
sulphur component, thereby disrupting the co-op- 229
erative role of the S-proteoglycans in the fibroblast 230
growth factor mediated signalling system that 231
maintains the overall growth and structural integ- 232
rity of the nervous system [21]. Disrupted S-pro- 233
teoglycan signalling systems are a consistent 234

235 feature of several neurodegenerative diseases,
236 such as TSEs [21–23]. Furthermore, the demon-
237 stration of a metabolic association between the S-
238 proteoglycan molecules and the cellular prion
239 protein [22,23] suggests that the disruption of S-
240 proteoglycan mediated signalling systems – ob-
241 served in the pathogenesis of TSEs – might perform
242 a pivotal role in the origins of TSEs.

243 **High intensities of low frequency acoustic**
244 **shock waves, a secondary prerequisite for**
245 **the pathogenesis of TSEs? (see Fig. 1)**

246 It is proposed that these Ag, Ba, Sr or Mn pollutants
247 act as founder nuclei which seed piezoelectric
248 crystals [29] that incorporate PrP and ferritin pro-
249 tein into their structure. Loss of S-proteoglycan
250 activity would deprive the biosystem of one of its
251 key endogenous molecules that inhibits the for-
252 mation of crystals; thereby enabling these rogue
253 crystals to multireplicate unimpaired, which
254 chokes up the formerly viable PrP-proteoglycan
255 conduits of electrical conduction throughout the

256 CNS. This compromises the ability of the brain to
257 process and protect itself against incoming bouts of
258 high energy sonic shockbursts from the external
259 environment – such as the intensive pressure
260 waves that radiate from sources of low frequency
261 infrasound – low flying jets, military explosions,
262 earthquakes, thunderstorms, etc. Interestingly,
263 one or other of these eco-phenomena have been
264 consistently observed as a key characteristic of
265 every global TSE cluster environment that has been
266 studied by the author to date [2,3].

267 **The piezoelectric crystal component captures**
268 **the sound**

269 Incoming mechanical shock waves of energy are
270 absorbed by the ferrimagnetically ordered ‘piezo-
271 electric’ crystals, and duly transduced into an
272 electrical/magneto energy [30] which largely ac-
273 cumulate within the crystalline metal-PrP-ferritin
274 aggregates [31] (e.g., the ‘prion fibrils’) until a
275 point of saturation polarization is reached.

276 In this respect, the whole TSE disease process
277 can be likened to a battery on continuous charge;
278 whereby the incoming energy from the environ-
279 ment is ultimately captured within these ferro-
280 electric crystal pollutants lodged in the brain.
281 Electric signals and Magnetic fields are generated
282 on the crystal surface. These upset the magneto-
283 electrical homeostasis of the CNS and initiate chain
284 reactions of deleterious free radical [32] mediated
285 spongiform neurodegeneration, leaving a ‘halo’ of
286 neuronal vacuolation around the crystal particles
287 (see Fig. 2). Interestingly, some of the barium
288 seeded crystals have a tendency to develop into
289 ‘flower’ shaped structures. In this respect, the
290 whole putative concept of the pathogenic crystal
291 as a cause of TSE could explain the presence of the
292 large ‘Florid plaques’ surrounded by a halo of
293 spongiform neurodegeneration – the neuropatho-

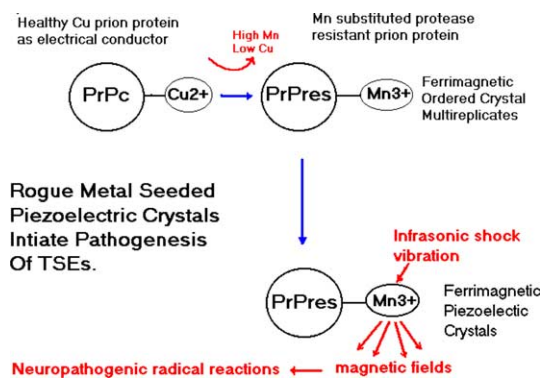


Figure 1

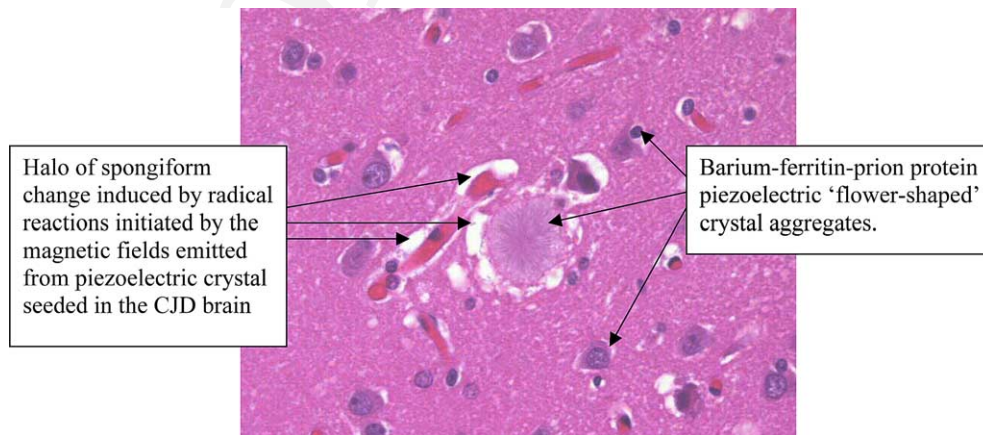


Figure 2

294 logical hallmarks of the brains of vCJD, Kuru and
295 CWD casualties (see Fig. 1).

296 The capacity of the metal based piezoelectric
297 crystal to transduce and store up incoming sound
298 energy is well illustrated in several industrial ap-
299 plications; such as the use of chromium, iron or Mn
300 3+ doped crystals in audio music tape material that
301 stores up a record of sound energy in ferromagnetic
302 form. Other examples involve the use of Ba in sound
303 proof boarding for recording studios, or in the mi-
304 crophone; where the incoming pressure waves of
305 sound are duly converted by the microphone's pi-
306 ezoelectric crystals into electrical signals.

307 The TSE diseased brain could be described as
308 having a million mini 'microphone' contaminants
309 lodged within it – yet without any loud speaker
310 system to dissipate the energy that has been
311 transduced from the sound. Each ferroelectric
312 crystal accumulates that energy until a point of
313 explosive saturation is reached.

314 **The additional effects of sonic shockwaves on** 315 **protein conformation**

316 Furthermore, the bombardment of the biological
317 system at large by shock waves from the external
318 environment will also invoke an intensive burst of
319 'molecular motion' in the tissues, where proteins,
320 for instance, are jiggled around by the heat energy
321 that results from the actual pressure of the in-
322 coming shock wave pulse. In more extreme cir-
323 cumstances, the protein molecules will adopt an
324 abnormal conformational shape as a result of the
325 pressure stimuli, but will rapidly refold back into
326 their normal shape providing those proteins are
327 attached to their correct metal co-partners. But in
328 the individual whose metallo proteins are conju-
329 gated onto rogue replacement metals – as is pos-
330 tulated in the case of the rogue metal
331 contaminated, TSE susceptible brain – any sonic
332 shock induced conformational derangement of
333 those proteins could remain as a permanent 'fix-
334 ture' – as evidenced by the presence of the stable
335 misfolded PrP isoform in TSE diseased brain [5].

336 Interestingly, PrPc cell cultures which have been
337 challenged by sound waves have demonstrated a
338 10-fold increased expression of PrP [33], This pro-
339 vides some evidence that PrPc expression responds
340 to incoming challenges of acoustic energy.

341 **Ag, Sr, Ba or Mn nucleated piezoelectric** 342 **crystals, the pathogenic transmissible agent** 343 **in TSEs?**

344 The theory of the proteinaceous prion particle as
345 the pathogenic agent in TSEs has not been vali-

dated in the experimental or epidemiological con- 346
text. It is proposed that the protein moiety of the 347
metallo-prion complex merely serves as an innoc- 348
uous *vehicle* that transports the toxic metal causal 349
agent; much like a trojan horse galloping around 350
the synapses of auditory, circadian circuits of the 351
brain carrying its lethal cargo of metallic missiles 352
on board – a magnetic field inducing 'fire power' 353
capacity that is potentially capable of detonating a 354
deleterious chain reaction of free radical mediated 355
neurodegeneration – a progressive pathogenesis 356
that can be generated by the magnetic fields and/ 357
or radioactive decay that are emitted by any rogue 358
magnetically ordered or radioactive metal piezo- 359
electric crystals that are successfully seeded in 360
biological tissue. 361

The magnetic fields proliferate, progressively 362
corrupting the key circadian/acoustic/vestibular 363
circuits (areas that are lesioned in TSEs [5,15,26]), 364
inducing a contagious domino-like aggregation of 365
metallo-crystal PrP molecules that multireplicate 366
themselves into crystalline 'fibril' tombstone de- 367
posits in the diseased brain. 368

Much of the epidemiological history surrounding 369
the major epidemic of BSE in the UK, indicates that 370
the protein only 'prion' hypothesis on the origins of 371
BSE fails to fulfil Koch's postulates [2]. For in- 372
stance, ≈10–25% of the cattle that have been 373
slaughtered each month under the UK govern- 374
ment's BSE order for exhibiting the full profile of 375
BSE symptoms had failed to demonstrate the 376
presence of prions at post-mortem [34]. The fact 377
that these so called 'BSE-negative' cows shared the 378
same idiosyncratic clinical profile and spatial- 379
temporal distribution as the BSE positive cows, 380
suggests that these *prion negative* cases were 381
suffering from the same disease as the *prion posi-* 382
tive cases. 383

Since transmission experiments using TSE af- 384
fected brain homogenate have indicated that the 385
causal agent remains 'pathogenic' after heating up 386
to temperatures in excess of 800° [35], then the 387
theory of Ag, Ba, Sr or Mn crystals as the TSE causal 388
candidate fulfils this prerequisite for pathogenicity 389
well. For the piezoelectric capacity of most ferro- 390
electric crystal structures will remain stable until 391
the crystals are heated up to their respective 'curie 392
point' temperatures around the 1000° mark; at 393
which point the orientation of the domains and the 394
alignment of the dipoles is destroyed, whereupon 395
the crystal is instantly depolarised and drained. At 396
higher temperatures, the melting point of the 397
crystal is exceeded [29]. Likewise, each ferrimag- 398
netically ordered metal atom within the crystal 399
structure would hold onto their remnant magnetic 400
charge until they are heated to temperatures above 401

402 their respective curie point temperatures
403 (500–600°) [8], whence the thermal agitation is
404 sufficient to instantly drain the charge. Such factors
405 as the piezoelectric and magnetic field inducing
406 capacity of the crystal, as well as its self replicating
407 properties, determine the pathogenic potential of
408 the crystal, thus its ability to induce TSE.

409 It is therefore proposed that these ferroelectric
410 crystal pollutants represent the transmissible, heat
411 resistant, pathogenic agents that cause TSE. These
412 crystals can be readily transmitted via any inocu-
413 lum of TSE contaminated brain homogenate into a
414 healthy laboratory animal, which, in turn, rein-
415 troduces the TSE crystal nucleating agent into the
416 healthy recipient organism.

417 **Methods**

418 **Soil sample collection/analysis method**

419 Each soil sample comprised a 300 g sample drawn
420 from a mix of 20 columns of dry soil bored with a
421 stainless steel auger; each column having been
422 bored at equidistant spaces along a W shape
423 spanning an area of ≈ 10 acres, the area being
424 representative of the region grazed by the CWD
425 affected deer under study. Each column was drawn
426 from the top soil to a depth of 6 in. having taken
427 care to avoid inclusion of root material/surface
428 organic matter and collection of samples near to
429 gateways, roadsides, animal dung, disturbed/ex-
430 cavated or polluted terrain. The 20 columns were
431 collected into a plastic bag, then mixed into an
432 even homogenate, from which a further sample of
433 no more than 300 g was drawn and placed into a
434 small polythene bag, then sealed, labelled and
435 transported to the laboratories at the Department
436 of Geology, Royal Holloway, University of London,
437 Egham Hill, Surrey TW20 0EX, where samples were
438 dried after arriving at the laboratory in forced air
439 flow cabinets. The temperature was maintained
440 below 32 °C during the 12 h drying period and the
441 air was constantly dehumidified. The soil samples
442 were then ground to pass a 2 mm mesh using a
443 hammer mill. The mill was flushed between sam-
444 ples using a small portion of the next sample. Each
445 sample was analysed by standard Mass Spectrom-
446 eter analytical procedure.

447 **Vegetation sample collection/analysis**

448 Each plant tissue sample comprised a 200 g sam-
449 ple representing tissue collected from ≈ 10 pick-

ings taken at equal spacings in a W shape across 450
an area of ≈ 10 acres that was representative of 451
the region grazed by the CWD affected deer/elk 452
under study. Samples were picked dry and at an 453
appreciable distance from roadsides, gateways, 454
animal manure, mechanically disturbed or 'spot' 455
polluted terrain. The tissue was packed directly 456
into plastic bags, lightly sealed, labelled, refrig- 457
erated and then transported to the laboratories of 458
the Department of Environmental Sciences at 459
Derby University, Kedleston Road, Derby, DE22 460
1GB, UK. Each sample was placed in a plastic 461
sieve and thoroughly washed in deionised water. 462
After removal of any roots or soil, the samples 463
were spread evenly on a drying tray and dried in a 464
90 °C oven to constant weight, and then ground by 465
Christy Norris mill, a small portion of the next 466
sample being used to flush the mill, before col- 467
lection of the ground material. The samples were 468
then prepared for analysis by dry ashing for non- 469
volatile elements and wet digestion in aqua/regia 470
for volatile elements (e.g., selenium). Analyses 471
was by standard ICP scan. 472

473 **Antler collection and analysis**

474 Antlers from 2 to 3 years old free ranging or 474
farmed cervidae were collected during April–June 475
2003 across the regions/farms where the most 476
intensive outbreaks of CWD had been officially 477
identified (DNR 2003). Samples were batched ac- 478
cording to CWD cluster/CWD-free region and sent 479
to the University of London at the Royal Holloway, 480
Egham, Surrey, UK for chemical analysis. The 481
samples for analysis were ignited to 600 °C to 482
remove organic material and then powdered in an 483
agate mortar and pestle. They were then dis- 484
solved in hydrofluoric and nitric acids and analy- 485
sed by Inductively Coupled Plasma Atomic 486
Emission and Inductively Coupled Plasma Mass 487
Spectrometry. 488

489 **Results and discussion**

490 **High silver, possible environmental sources** 490 491 **and modes of uptake**

492 High levels of Ag and low levels of Cu were re- 492
corded in the antler material, soils and deer 493
browse vegetation drawn from CWD affected zones 494
(see Tables 1 and 2, Graphs 1 and 2). These results 495
represent the first time that Ag has been detected 496

Table 1 Levels of metals in antler from CWD cluster and CWD-free zones across North America

Matrix	Sampling zone	CWD status	Ca0 (%)	Ba (ppm)	Cu (ppm)	Sr (ppm)	Ag (ppm)	Mn (ppm)
Antler	Fort Collins, Co.	CWD+	25.25	138	2	130	3.9	0 (w)(t)
Antler	Fort Collins, Co.	CWD+	25.56	125	2	137	3.4	0 (w)(b)
Antler	Mt Horeb, Wi	CWD+	25.78	63	3	42	2.1	0 (w)(t)
Antler	Mt Horeb, Wi	CWD+	25.43	60	3	42	3.0	0 (w)(b)
Antler	N.Manitou, Sk	CWD+	25.59	206	2	117	4.7	0 (w)
Antler	N. Manitou, Sk	CWD+	25.64	202	2	130	2.7	0 (w)
Antler	N. Manitou, Sk	CWD+	25.29	280	2	114	9.6	0 (w)
Antler	Lloydminster, Sk	CWD+	24.98	88	76	136	3.1	0 (F)elk
Antler	Lloydminster, Sk	CWD+	25.20	77	3	120	2.7	0 (F)elk
Antler	Lloydminster, Sk	CWD+	25.35	156	2	130	4.6	0 (F)elk
Antler	Manitou, Sk	CWD+	24.17	184	2	121	0.7	4 (w)
Antler	Manitou, Sk	CWD+	23.96	179	3	122	0.9	4 (w)
Antler	Manitou, Sk	CWD+	23.98	253	3	140	3.1	8 (w)
Antler	Manitou, Sk	CWD+	24.12	258	3	141	1.0	8 (w)
Antler	Manitou, Sk	CWD+	23.29	269	4	145	0.8	8 (w)
Antler	Manitou, Sk	CWD+	24.28	263	3	143	9.6	8 (w)
Antler	Manitou, Sk	CWD+	23.71	185	3	122	0.8	4 (w)
Antler	Manitou, Sk	CWD+	24.04	191	3	124	5.2	4 (w)
Antler	Manitou, Sk	CWD+	24.32	197	2	115	2.8	8 (w)
Antler	Manitou, Sk	CWD+	24.27	200	2	117	2.8	8 (w)
Antler	Manitou, Sk	CWD+	24.50	202	2	116	2.5	8 (w)
Antler	Manitou, Sk	CWD+	24.59	202	2	117	0.8	4 (w)
Mean CWD antler			24.78	181	5.8	119	3.2	3.4
Antler	Alberta	CWD-free	25.98	56	2	77	2.4	0 (F)(t)
Antler	Alberta	CWD-free	25.35	52	2	71	4.2	0 (F)(b)
Antler	Alberta	CWD-free	25.10	72	2	38	2.0	0 (w)(t)
Antler	Alberta	CWD-free	24.73	69	3	36	3.4	0 (w)(b)
Antler	Somerset UK	CWD-free	24.15	38	3	42	0.0	8 (w)
Antler	Somerset UK	CWD-free	24.57	43	3	57	0.1	19 (w)
Antler	Somerset UK	CWD-free	23.84	74	3	45	0.1	12 (w)
Antler	Somerset UK	CWD-free	24.08	66	2	47	0.0	23 (w)
Antler	Somerset UK	CWD-free	24.20	42	2	42	0.0	4 (w)
Antler	Somerset UK	CWD-free	24.18	43	3	42	0.0	43 (w)
Mean CWD-free antler			24.61	55	2.5	49	1.2	11
Reference mammalian bone			25	5	13	52	0.01	0.2

(w) = Antler from wild deer herd, (F) = antler from farmed deer herd, (t) = section from tip of antler, (b) = section from base of antler.

497 in antlers, whilst adding some support to the pro-
498 posal [2,3] that high Ag and low Cu in the envi-
499 ronment may bring about an Ag replacement of
500 vacant Cu ligands on the cellular prion protein
501 (PrPc).

502 Particularly interesting is the fact that Ag was
503 virtually undetectable in antlers collected from
504 CWD-free herds in the UK. However, antlers col-
505 lected in the CWD free areas of Alberta (which
506 adjoins the CWD cluster zones along the Alberta/
507 Saskatchewan borders) revealed marginally ele-
508 vated levels of Ag, which may indicate that these
509 areas are approaching the threshold of 'high risk'
510 for hosting outbreaks of clinical CWD in their deer
511 populations in future.

Ag is potentially highly toxic [36], exerting a
strong competitive binding affinity for specific Cu
ligands on cuproproteins [37]. The degree of in-
toxication encountered following Ag exposure is
controlled by the overall Ag/Cu ratio within the
biosystem.

512 Apart from the naturally occurring sources of Ag
513 in soils – well renowned in the Colorado CWD
514 cluster area – possible routes of Ag exposure in the
515 CWD cluster ecosystems could stem from routine
516 feeding of Ag contaminated concentrated feed
517 pellets to captive and wild deer herds. In this re-
518 spect, Ag was measured at 2.2 ppm in the feed
519 samples collected from deer farms across North
520 America during this study (see Table 1).
521
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523
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526

Table 2 Levels of metals in soils and vegetation sampled across CWD cluster and CWD-free zones

Matrix	Sampling zone	CWD status	CaO (%)	Ba (ppm)	Cu (ppm)	Sr (ppm)	Ag (ppm)	Mn (ppm)	S (%)
Soil	Colorado	CWD+	2.65	568	18	192	0.35	619	0.27 (40)
Soil	Wisconsin	CWD+	1.28	477	16	114	0.21	915	0.13 (40)
Soil	Saskatchewan	CWD+	1.14	905	24	193	0.27	853	NR (8)
Soil	Mean	CWD+	1.69	650	19	166	0.27	795	0.20
Soil	Vermont	CWD-	1.99	474	22	98	0.08	757	NR (20)
Soil	Alberta	CWD-	3.21	537	17	124	0.30	550	NR (17)
Soil	Mean	CWD-	2.60	505	19	111	0.19	653	
Soil mean reference			1.00	250	30	80	0.07	750	0.30
Veg	Colorado	CWD+	10,192	56	13	61.4	0.459	196	0.19 (40)
Veg	Wisconsin	CWD+	10,288	56	16	57.0	0.858	122	0.30 (40)
Veg	Saskatchewan	CWD+	11,295	50	4	68.1	NR	60	NR (6)
Veg	Mean	CWD+	10,590	54	11	62.1	0.658	126	0.24
Veg	Vermont	CWD-	7400	24	25	16.2	0.242	111	0.47 (20)
Veg	Alberta	CWD-	6271	31	5	16.4	NR	102	NR (2)
Veg	Mean	CWD-	6835	27	15	16.3	0.242	106	0.47
Pasture mean reference			5000	10	20	20	0.05	50	0.35
Concentrated feed pellets			1.83	16	55	25	2.2	222	

Analyses was performed by MS. Measurements relate to total levels of element recorded as ppm on dry basis. (20) = number of sample sites (covering approx 10 acres for each site) involved in the constitution of each mean level of metal displayed above. NR = not recorded.

527 Ag use in cloud seeding weather modification

528 Another significant source of Ag contamination in
529 the drought prone regions where CWD has
530 emerged, stems from the extensive aerial spray
531 application of silver iodide crystals used as foun-
532 der nuclei in cloud seeding 'rainmaking/snow-
533 making' operations [38]. The resulting Ag
534 contaminated rain permeates the local vegeta-
535 tion, as well as the growing crops that are in-
536 corporated into the concentrated feeds for the
537 deer.

538 Whilst airborne Ag can be absorbed directly
539 into the brain via the nasal-olfactory route of in-
540 halation [39], the 'fall out' from atmospheric Ag
541 contamination will bio-accumulate in bryophytes
542 (lichens/mosses concentrate Ag up to 9 ppm [38]
543 and other vegetation) (see Table 2), which are
544 subsequently ingested by the local deer/elk pop-
545 ulations.

546 It is interesting that the practise of cloud seed-
547 ing is largely contained within the North American
548 continent – the area which has hosted virtually all
549 cases of TSE in wild animals – whereas the appli-
550 cation of silver ions as a broad spectrum biocide
551 [40] in food production, etc., has been viewed with
552 greater caution by the US authorities.

Ag use as a biocide

There is a greater use of Ag for its biocidal poten-
tial within Europe; where it has been increasingly
used over the last two decades as a water purifier

553

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ENVIRONMENTAL PREREQUISITES UNDERPINNING UK's BSE EPIDEMIC;

1. Compulsory high dose use of systemic organo dithiophosphate insecticides;

*a. chelates copper, starving prion protein of its
copper co partner.*

*b. Increases permeability of b/b/brain
barrier – increasing uptake of rogue metal
crystal founder nuclei into the brain*

2. Exposure to Ba, Ag, Mn, Sr crystal nuclei;

* *Ba in fishmeal derived from North sea ?*

* *Ag in MBM due to use as salmonella
biocide in poultry units ?*

* *Sr90 from Chernobyl fall out ?*

* *Mn as high dose mineral inclusions?*

3. Exposure to Low Frequency shock waves;

* *Low flying military jet practise, Concorde
and other civilian jet take off overflights ,etc.*

Figure 3

557 and sterilising agent in establishments like the
558 London zoo, Rendering plants, Hospitals (for ster-
559 ilising surgical implements, etc.), dentists (also
560 used as a component of amalgam fillings), catering
561 establishments, dairy farms, etc. Establishments
562 which have been associated with high incidences of
563 TSEs. Ag biocides are also used in air conditioning,
564 waste water treatment, aquaculture, food and
565 beverage treatment, swimming pools and surface
566 cleaning in many applications.

567 It is interesting that the use of Ag ions to curb
568 salmonella escalated in the UK poultry industry
569 after the 'Edwina Curry' salmonella crisis hit UK
570 poultry farms in 1988; and the subsequent bioac-
571 cumulation of Ag through the farm animal food
572 chain (via use of waste poultry meat and bone meal
573 and manure as both a feed and fertiliser) could
574 have been contributory to the UK's BSE epidemic

that peaked in 1992 [34] (see Fig. 3) – where si-
575 multaneous exposure to the exclusive compulsory
576 high dose use of the Cu chelating organo dithio-
577 phosphate for warble control of UK cattle [1] had
578 deprived PrPc of its Cu co-partner, rendering the
579 protein vulnerable to an Ag replacement.
580

581 High incidence clustering of BSE has consistently
582 existed amongst cattle pastured in the main poul-
583 try/turkey producing region of Norfolk since BSE
584 first erupted [34], where surrounding farmland has
585 been generously fertilised by Ag and Mn rich poul-
586 try manure for many years.

High barium and strontium, possible environmental sources and routes of uptake

587
588
589 The other unusual observation resulting from this
590 study implicates the elevation of Ba/Sr and low

<i>Location</i>	<i>State</i>	<i>TSE type</i>	<i>Military connection</i>	<i>sonic source</i>	<i>Ba</i>	<i>Sr</i>	<i>Ag</i>	<i>Cu</i>
Tucson	AZ	CJD cluster	Missile factory staff	?				
Fort Collins	CO	CWD cluster Wild/captive deer.	Missile silos/ LF jets Rocky Flats nuclear weapons factory.	Quarry explosions Guns / LFjets Tectonic rift line	56	61	.46	13
Mt Horeb*	WI	CWD cluster	Hercules flightpath LF jets / Badger Munitions site	New Road blasting Guns / LF jets Quake epicentre	56	57	.85	16
White Sands * Missile Range	NM	CWD cluster	Missile test range	Missile explosions	83	35	1.42	10
Mission	TX	Scrapie cluster	Former Airbase (now private)	Under take off flight path			not sampled	
Allenstown	PA	CJD cluster	International Airbase	LF jets.			not sampled	
Garden State	NJ	CJD cluster	Fort Dix military camp MacGuire Airbase	LF jets / Guns Shell explosions			not sampled	
Mabton * Spokane *	WA WA	1 st BSE 1 st vCJD	Hanford Nuclear Pl Yakima Military train Camp / Othello Airbase	LF jets Shell explosions			not sampled	
Namao *	AL	Ist CWD farmed deer	Namao Airbase	Under take off flight path	61	40		9
Leduc *	AL	US BSE cow reared.	Leduc International Airport- mainly civilian	Under take off flight path	83	72		8
Tulliby Lake *	AL	Ist BSE	Cold Lake airbase Hercules flight path	Under LF jet practise circuit / Hercules flight path	45	100		4
Hillmond *	SA	CWD cluster farmed elk	Hercules flight path	Under Lloydminster airport take off path Gas well pumping	55	98		10
Manitou *	SA	CWD cluster wild deer	Camp Wainwright tank range	Tank shelling. Shooting range	64	53		3
Lloydminster *	SA	Ist vCJD	midway Cold Lake/ Camp Wainwright	Under Hercules flight path / tank shelling.	50	41		1.8

Figure 4 Location of the most renowned long standing and recent TSE clusters/outbreaks in USA/Canada in respect of their metal profile and proximity to major sources of low frequency sonic shock waves. LF = low flight. * = TSE case/s emerged since 2000.

591 levels of sulphur in the antlers, vegetation and soils
592 of the CWD affected deer (see Tables 1 and 2,
593 Graphs 1 and 2).

594 Since an insufficient number of studies have
595 been conducted on the levels of these metals in
596 antlers [41,42], the mean reference levels of Ba/Sr
597 in bone material have been used in this report as
598 the best alternative for providing mean reference
599 ranges of Ba/Sr in antlers. In respect of the refer-
600 ence levels of 5 ppm Ba and 52 ppm Sr in bone
601 matrix [43–45], the mean levels of 181 ppm Ba and
602 119 ppm Sr recorded in the CWD antlers in this
603 study could be regarded as 'elevated'. Levels of 55
604 ppm Ba and 49 ppm Sr in CWD-free antlers col-
605 lected in both the UK and Alberta were consistently
606 more than 3× and 2.5× lower than mean Ba/Sr
607 levels in CWD+ antlers.

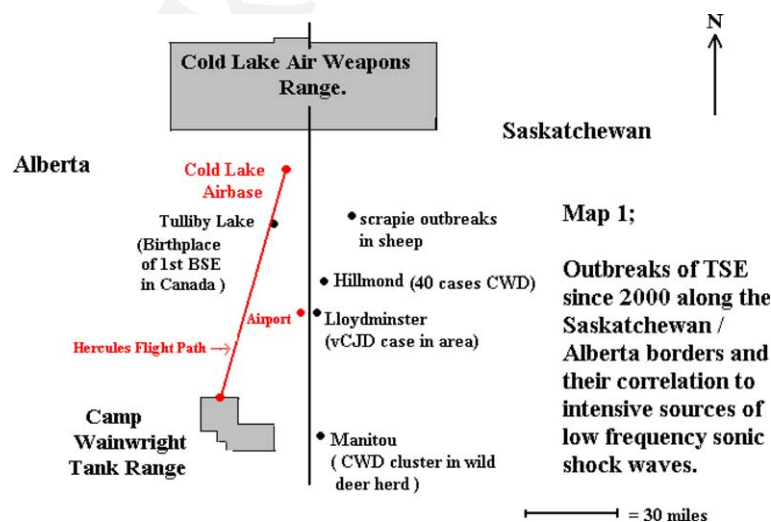
608 The mean levels of Ba and Sr were significantly
609 higher in the vegetation of the CWD cluster envi-
610 ronments at 54 ppm Ba and 62.1 ppm Sr in relation
611 to levels of 27.5 Ba and 16.3 Sr recorded in CWD-
612 free control areas. The levels in CWD clusters were
613 also several fold higher than their mean reference
614 levels of 10 ppm Ba and 20 ppm Sr [44,46] for
615 vegetation. A high Ba/Sr and low sulphur mineral
616 profile has also been recorded by the author in TSE
617 cluster ecosystems in Southern Italy, Sardinia, Ja-
618 pan and Iceland.

619 Sources of radioactive Sr90 in TSE clusters

620 Raised levels of Sr have been recorded previously in
621 a study on antlers [47] where exposure to atmo-
622 spheric contamination by radioactive Sr90 was
623 considered to be responsible. Radioactive counts
624 are currently being executed on the antler material
625 collected in this study, but are not yet complete.

If the high Sr (119 ppm mean) recorded in the
antlers in this study turns out to stem from a ra-
dioactive Sr90 source, rather than the stable Sr 88
form, then the contamination of the Northern
Hemisphere by Sr90 as a 'fall out' legacy of the
1986 Chernobyl accident and the 1960s/1970s nu-
clear weapons testing [48,49] could provide an
explanation for the source of the contamination.
Furthermore, the fall out from a more local source
involved the long term atmospheric leak of radio-
active metals from the Rocky Flats Nuclear Weap-
ons Plant during 1967 [50,51]. This was
environmentally monitored and found to contami-
nate [53,54] the precise region where the first re-
corded cluster of CWD emerged in north eastern
Colorado in 1968 [52]. The first deer to develop
CWD in this outbreak [52] had actually originated
from the same pens at the Fort Collins wildlife fa-
cility that had been involved in a raft of experi-
ments to monitor the effects of exposure to various
radioactive metals (Pu, Cs137, Sr90), which in-
cluded transporting and grazing the deer on the
intensively contaminated pastures around the
Rocky Flats Plant itself [53–55].

In this respect, it is interesting that the majority
of TSE clusters in North America have emerged
near to significant military munitions production/
storage/testing facilities such as the White Sands
missile range [3], a missile factory in Tucson [57],
The Rocky Flats Nuclear weapons factory [58], a
battery of 'cold war' missile silos scattered be-
tween NE Colorado/SE Wyoming, and the Cold Lake
Air Weapons Range/Camp Wainwright on the Al-
berta–Saskatchewan borders [58] (see Fig. 4, Map
1) where radioactive metal based materials are
known to have been used.



Map 1

662 Sr90 could represent a rogue metal candidate
663 that potentially initiated the intensive outbreak of
664 BSE across NW Europe in November 1986 – due to
665 the fall out of this metal in the rainstorms which
666 immediately followed the Chernobyl nuclear reac-
667 tor accident in April 1986 [48]. This could be rele-
668 vant to the many anecdotal reports by UK vets and
669 farmers that cite a prevalence of osteoporotic-like
670 bone wastage conditions and protracted episodes
671 of atypical hypocalcaemia ‘milk fever’ (that failed
672 to respond to standard therapeutic doses of Ca) in
673 cattle that later went on to develop BSE [59]. This
674 could indicate a case of successful Sr or Ba sub-
675 stitution at Ca binding domains [43] throughout the
676 biosystem.

677 Ferritin protein is an integral component of
678 the prion fibril [31] as well as acting as the
679 specific chelating agent for treatment of Sr90
680 poisoned mammals [56]. If radioactive Sr90
681 binding to PrP/proteoglycan/ferritin could be
682 experimentally achieved in cell culture models,
683 then it would be tempting to postulate a causal
684 association between chronic exposure to the
685 more reactive Sr90 species and the more short
686 lived, aggressive ‘new strains’ of TSE; where the
687 additional pathogenic complication of radioac-
688 tive decay emitted from the Sr90–PrP-ferritin
689 crystalline complexes would exacerbate the in-
690 tensity of neuropathogenic free radical chain
691 reactions, creating an overall acceleration of the
692 standard duration of the clinical phase of the
693 conventional TSEs. This would invoke clinical
694 disease in younger as opposed to more elderly
695 mammals.

696 **Use of Ba, Sr and Ag in conventional munitions**

697 The toxic common denominator that underlies
698 this correlation between the close proximity of
699 military bases to TSE clusters, may actually re-
700 late to the contamination of their surrounding
701 environments with the more conventional, non-
702 radioactive metals that stem from the use of Ba,
703 Sr or Ag in military munitions and other applica-
704 tions.

705 Furthermore, the previously reported ‘sonic
706 shock’ prerequisite that has been observed in every
707 significant global TSE cluster visited by Purdey [2,3]
708 is also evident at the majority of the military in-
709 stallations that are contiguous to these TSE clusters
710 in North America [2,3] (see Map 1).

711 For example, many of these TSE affected animal/
712 human populations had been found to be living
713 beneath low fly jet flight paths or the ‘take off’
714 flight paths coming out of military or civilian air-
715 bases like Namao and Leduc on the north and south
716 sides of Edmonton, respectively.

Use of Ba ions in aerosol applications employed by the military and geophysical researchers

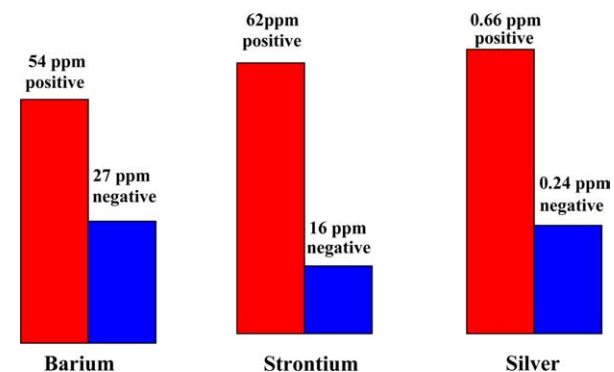
719 It should also be noted that aerosols containing the
720 Ba ion – such as the ferroelectric Ba stearate or Ba
721 strontium titanate compounds [60,61] – are dis-
722 charged along jet flight paths/low flight practise
723 areas and around munitions production/storage/
724 guided missile testing facilities as a means of en-
725 hancing/refracting radio/radar signalling commu-
726 nications for maintaining a reliable measure of
727 security and rapid communication around the cur-
728 vatures of the earth. Ba ions have also been widely
729 discharged into the atmosphere since the mid
730 1970s as a means of conducting geophysical ex-
731 plorations of the ionosphere [62].

732 Thus any foodchain that is sited around these
733 top security military locations, flight paths or be-
734 neath these areas of ionospheric exploration, could
735 find itself subjected to the toxic ‘fall out’ from this
736 mode of atmospheric metal pollution.

Naturally occurring geochemical and bioconcentrated sources of Ba/Sr

739 The elevated levels of Ba in the North American
740 CWD clusters may also partly derive from the do-
741 lomite/limestone and Cambrian granitic mica
742 schist soil types of the CWD cluster areas [43].
743 These light, low organic matter soil types are nat-
744 urally high in Ba and Sr [43], whilst being notori-
745 ously low in sulphur and copper. The low sulphur
746 perspective exacerbates the problem of Ba/Sr
747 toxicity in the mammal who is dependent upon
748 these foodchains, in that an available source of
749 free sulphur in the soil will conjugate with Ba and
750 Sr, thereby locking up those minerals and acting as
751 a ‘toxic sink’/preventative against Ba intoxication
752 [45].

753 The customary spreading of spent barium drilling
754 mud across farmland (a waste product of the fast

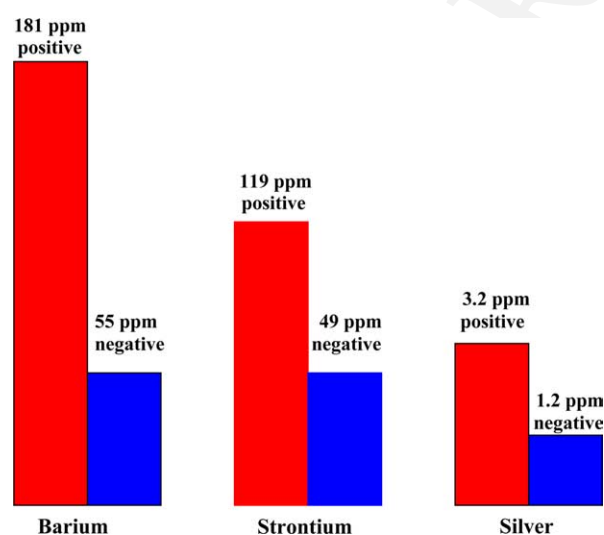


Graph 1 Comparative levels of metals in the pasture of CWD and CWD-free regions. (Mean averages drawn from 86 and 22 samples, respectively.)

755 expanding oil and gas well industry in the CWD
756 areas) has compounded the problem; with sub-
757 sequent uptake of Ba into the pasture and hay
758 crops which are ingested by local cervidae popu-
759 lations. Cultivated plants such as alfalfa/soy bean,
760 as well as the wild 'locoweed' flora are prevalent in
761 the CWD areas and are also ingested by the cervi-
762 dae populations. These species are renowned to
763 bioconcentrate Ba and Sr to high levels [43,44,63].

764 Drought conditions exacerbate the uptake of Ba/ 765 Sr into the ruminant

766 The author has observed that conditions of pro-
767 longed drought precede the outbreaks of CWD in
768 North America. This correlation could be linked to
769 several eco-influences that surface during drought
770 conditions; such as the aforementioned use of Ag/
771 Ba as cloud seeding nuclei during dry seasons.
772 Drought conditions also exacerbate the problem of
773 metal bioconcentration in grazing deer, in that the
774 resulting shortages of pasture cause malnourished
775 cervidae populations to consume abnormally high
776 intakes of pine, juniper and locoweed as a sub-
777 stitute for their normal rations – a phenomena that
778 is widely reported by hunters and ranchers oper-
779 ating in the CWD environments. These plants bio-
780 concentrate Ba and Sr [43]. Furthermore, when
781 overpopulated deer are forced to compete for the
782 dwindling reserves of close cropped, drought-par-
783 ched pasture, their intake of topsoil – and the
784 metals contained therein – is dramatically in-
785 creased [51,64]. Interestingly, increased amounts
786 of soil, sand and grit have been observed in the
787 digestive tracts of CWD positive slaughtered deer in
788 relation to CWD-free deer [52].



Graph 2 Comparative levels of metals in the antlers of deer from CWD and CWD-free regions. (Mean averages drawn from 22 and 10 antler samples, respectively.)

It should also be noted that some of the con-
centrated feeds that were fed on deer farms during
the drought periods contained Ba and other crystal
nuclei in the 'bentonite' clay material that is ad-
ded to the feed pellet as a binder. Bentonite was
also used at a high inclusion rate in cattle con-
centrated feeds manufactured in the UK, where
BSE erupted at epidemic levels.

Conclusion and future research

Irrespective of any role that elevated levels of Ag,
Ba and Sr may play in the pathogenesis of TSEs, the
novel observation of Ag in antler might suggest that
the antler acts as a hitherto unrecognised toxic
'sink' for storing excess intakes of Ag, Ba, Sr and
other metals in cervidae who are thriving off
foodchains that have bioconcentrated these met-
als. The toxic load is conveniently shed along with
the antler on an annual basis. Or, alternatively, Ag
and/or Ba could perform some metabolic role as an
electrical superconductor for mediating the rapid
growth of the antler.

Whilst previous field studies had observed high
levels of Mn and low levels of Cu in Icelandic and
Slovakian TSE cluster environments [1,2], these
more recent studies – which analysed for a broader
spectrum of metals – have found elevated Ag/Ba/
Sr and low Cu in the North American TSE clusters.
These latest advancements indicate that the lab
experiments conducted by Dr. David Brown [65] –
which successfully generated 'de novo' protease
resistant PrP formation after introducing Mn into
Cu deprived PrP cell cultures – need to be ex-
tended in order to test the impact of these addi-
tional cations in the Cu depleted PrP/proteoglycan
cell culture models. The rogue metal loaded cells
should be exposed to low frequency sonic shock
waves as a second stage challenge. Furthermore, It
would be interesting to run cell culture tests to see
if any radioactive metals, like Sr90, will bond onto
PrPc/proteoglycans in the absence of Cu/Zn.

These environmental observations indicate that
the high risk TSE foodchains need to be fortified
with copper and/or zinc additives, administered in
feed supplement or fertiliser form. These could
putatively act as a preventative against the emer-
gence of TSEs in susceptible individuals. Such
measures would guarantee an optimum dietary in-
take of Cu and Zn, enabling adequate Cu/Zn bind-
ing to PrPc's octapeptide metallo-domains or
proteoglycan centres, thereby protecting these
molecules against competitive substitution by ro-
gue replacement metals, that could act as founder

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841 nuclei for seeding crystals and initiating the path-
842 ogenesis of TSE.

843 The therapeutic use of compounds which can
844 cross the blood brain barrier and inhibit the prop-
845 agation of metal nucleated crystals by chelating
846 sulphate, silicate, etc., could be tested as a means
847 of arresting clinical TSE [66]. In this respect, it is
848 interesting that anti-malarial quinine molecules,
849 which have been shown to arrest the development
850 of TSEs, are also recognised to chelate alkali earth
851 metals, like Ba and Sr [67].

852 Since environmental exposures to the soluble
853 sources of Ba will chelate sulphur in the biosystem
854 [43] – thereby disrupting the viable synthesis of
855 the sulphur dependent proteoglycan molecules – it
856 is interesting that the therapeutic use of the S-
857 proteoglycan heparin molecules is having a positive
858 effect in human victims of vCJD. These beneficial
859 effects could merely result from the reintroduction
860 of viable sulphated proteoglycan molecules into
861 the proteoglycan depleted biosystem; where the
862 therapeutic arrest of the progression of the disease
863 could be linked to the reactivation of proteoglycan
864 mediated growth factor signalling, combined with
865 the ability of the restored proteoglycans to inhibit
866 any further multireplication of crystals.

867 More extensive and detailed analytical studies
868 need to be performed on these environmental
869 perspectives of TSEs in order to reach a more
870 concise and conclusive consensus on these pre-
871 liminary observations within this interesting area of
872 TSE research.

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884 and the biophysical effects of acoustic pressure
885 waves, respectively.

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