Case Report

Mercury Vapour Long-Lasting Exposure:
Lymphocyte Muscarinic Receptors as Neurochemical
Markers of Accidental Intoxication

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1. Introduction

Prolonged exposure to mercury (Hg) vapour may result in clinical pictures of chronic poisoning. This lasting intoxication, usually due to Hg vapour inhalation, is characterized by an initial flu-like syndrome, affecting at the outset the respiratory tract with symptoms of cough, sore throat, shortness of breath, and chest pain, then followed by signs affecting gastrointestinal, central, and peripheral nervous system, with a wide range of symptoms including fever, erythematous rash, itching, chills, gastrointestinal complaints, metallic taste, headache, and weakness. Anyway this “metal fume fever,” a syndrome commonly confused with a viral etiology, still remains a poorly understood syndromic picture [1].

Mercury poisoning is mainly described and reported in medical literature as a result of occupational exposure. Nonetheless, chronic exposure to Hg vapour is also possible in domestic/nonoccupational setting. Particularly, with the wide utilization of Hg in thermometers, sphygmomanometers, and barometers, mainly used at home and in hospitals as well as in schools, an accidental breakage of these devices may cause spillage of Hg droplets resulting in a chronic elemental mercury intoxication [1–3].

Additionally, indoor unintentional exposure may involve domestic appliances, such as the newest artificial lighting systems, for example, compact fluorescent lamps (CFLs), which are known to be energy-efficient compared to incandescent bulbs but contain milligram (mg) quantities of...
Hg. Particularly, international concerns have been raised regarding potential Hg vapour exposures following CFLs breakage, and various efforts have focused on managing this issue [4, 5].

Mercury used in consumer products is metallic Hg. Inhalation is usually the main route of concern because 80% of inhaled Hg is absorbed. After inhalation, elemental-Hg is readily absorbed through the alveolar membrane and transported by the blood to the brain and other target tissues. The susceptibility of central nervous system (CNS) to Hg is well established according to epidemiological and experimental investigations. Moreover, substantial evidences showed that cholinergic muscarinic system can be affected by in vitro and in vivo exposure to Hg [6]. Indeed literature data demonstrated that some environmental neurotoxic chemicals, other than Hg, may influence cholinergic muscarinic system by a variety of mechanisms. For example, organophosphates interact directly with receptor protein, acting either as agonist or as antagonist. Moreover, other agents may alter the receptors indirectly, either by changing the levels of endogenous neurotransmitter acetylcholine (as in the case of organophosphorus insecticides) or by damaging muscarinic receptor-bearing cells (e.g., trimethyltin) [7].

Concerning the Hg, because the body eliminates this metal slowly, cumulative exposure is the primary matter of concern, being the cause of a wide range of heavy health adverse effects [8].

Mercury chronic poisoning syndrome includes neuropsychiatric disturbances as well as peripheral neuropathy and renal involvement (presenting as proteinuria or tubulopathy). In particular, neurological symptoms may include decreased nerve impulse conduction, decreased motor skills (e.g., finger tapping, and hand-eye coordination), irritability, poor concentration, shyness, tremors (initially affecting the hands and sometimes spreading to other parts of the body), incoordination (e.g., difficulty walking), and short-term memory loss. The motor skill effects may be reversible, but short-term memory loss may be permanent [8]. Moreover, severe hypertension due to catecholamine excess was described in previous reports [2].

From a clinical point of view, misdiagnosis of Hg poisoning, often as a flu-like syndrome (at early onset) or as a psychological disorder (at later stages), is a common problem. Sometimes, before the correct diagnoses, patients worsen after returning to the place of contamination [8].

Because exposure to neurotoxics, including Hg, may cause biochemical and molecular events indicating early-stage effects of exposure preceding the onset of overt disease, monitoring these early events may represent a valuable approach, employing neurotoxicity markers as useful tool for detecting subclinical disease states and initial detrimental changes associated with long term low-dose exposure to Hg, thus supporting the clinicians in making an early differential diagnosis.

We report a case of chronic, nonoccupational mercury poisoning due to 10-year prolonged Hg vapour exposure as a result of spillage from broken barometers at home, parallelled by related alteration in peripheral neurochemical parameters, that is, lymphocytes muscarinic receptors. Our laboratory data clearly supported the use of this peripheral biomarker as susceptible target for Hg neurotoxicity in human.

2. Case Presentation

A 72-year-old man (70 kg body-weight) presented to our Toxicology Unit (TU) with a 10-year past medical history of progressive neurological symptoms (Table 1), to investigate a polyneuropathy of suspected toxic origin. The patient's occupational history was negative for previous exposure to metals. A nuclear magnetic resonance (NMR) performed 5 years before was negative for brain lesions. The patient mentioned the presence of a big broken barometer at his home, maintained near heating source in his study-room during the last ten years.

Notably, in October 2010, after a private consultation with a Belgian neurologist in Anversa, abnormal enhanced BHg levels were determined, that is, 36 microg/L at first control. Thus, chelation therapy cycles with administration of 2,3-dimercapto-1-propanesulfonic (DMPS) acid were prescribed by the Belgian physician. This therapy scarcely diminished BHg levels (26.7 and 21 microg/L after the first and second chelation cycles, resp.), also failing to produce any healthy relief on neurological symptoms (i.e., postural instability and sensory-motor polyneuropathy lasting) (Table 1). We may suppose that therapy inefficacy could be related to underdose, incorrect administration, and scarce adherence to the therapy, although these hypotheses are speculative since the patient was not directly managed by our Toxicology Unit during DMPS treatment. Moreover, it has to be mentioned that DMPS administration may be associated with side effects, such as allergic reactions and lowering in blood pressure, which could have contributed to a decrease in an elderly patient's compliance.

Hence, the patient was hospitalized (June 2011) at our clinical Toxicology Unit of IRCCS Salvatore Maugeri Foundation (FSM), Scientific Institute of Pavia. Toxicological clinical evaluation confirmed the altered neurological picture documented across the last decade (Table 1), characterized by motor ataxia, postural instability, positive Romberg, increased motor tone, paresthesias, and sensory deficits in-touch at inferior limbs. Neurophysiological tests (e.g., motor-sensory electroneurography (ENeG) and vegetative nervous system compartment electroneurography) revealed a mild axonal sensory-motor polyneuropathy at superior and inferior limbs. Parallel neuropsychological evaluation (NPE) demonstrated a normal global cognitive functioning. A standard electroencephalogram (EEG) was read as within normal limits.

Standard laboratory investigations demonstrated mild anisopikilocytosis, microcytosis, and anemia, consistent with typical heterozygous beta-thalassemic traits. Furthermore, specific haematochemical tests (Table 2), revealed an increased concentration (i.e., 67 microg/dL versus normal value: < 40 microg/dL) of erythrocyte zinc protoporphyrin (ZnPp) only. In accordance with previous clinical investigations [9, 10], we related this latter slightly altered parameter to the patient thalassemic phenotype, since the value measured
Table 1: Clinical and neurological events/evaluations during the decade (2000–2011) before admission to our Toxicology Unit.

<table>
<thead>
<tr>
<th>Year</th>
<th>Event/Consultation Details</th>
<th>Medical neurological division/consultant</th>
<th>Lab test/analyses</th>
<th>Symptoms/diagnosis</th>
<th>Therapy</th>
</tr>
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<tbody>
<tr>
<td>2000–2005</td>
<td>Neurological alterations onset and progression: postural instability during deambulation, associated with paresthesias and hypoesthesia at anterolateral surface of thighs; in 2003, after prostatectomy, hypoesthesia extended to tailbone area, paralleled by pain worsening at inferior limbs</td>
<td>San Raffaele Hospital, Cefalù, Italy</td>
<td>(i) Supra-aortic trunks and inferior limbs color-Doppler (CD)</td>
<td>(i) Sensory-motor polyneuropathy at inferior limbs</td>
<td>Gabapentin: lack of detailed pharmacological plan documentation</td>
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<td>(ii) Electromyography (EMG) and motor evoked potentials (MEP)</td>
<td>(ii) Hyperintense Punctate frontobilateral subcortical foci of gliosis</td>
<td>Therapeutic drug treatment self-suspended by the patient</td>
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<td>(iii) Encephalic and spinal column magnetic resonance imaging (MRI)</td>
<td>(iii) Spinal disc herniations and lumbar discal bulging</td>
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<td></td>
<td>(iv) Vertebral hemangiomas in some metameric segments at dorsal and lumbar levels</td>
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<td>2006</td>
<td>(hospitalized)</td>
<td>San Raffaele Hospital, Cefalù, Italy</td>
<td>(i) Electromyography (EMG), motor evoked potentials (MEP), somatosensory evoked potential (SSEP) monitoring</td>
<td>(i) Sensory-motor polyneuropathy</td>
<td>Lack of documentation</td>
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<td></td>
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<td>(ii) Sural nerve biopsy</td>
<td>(ii) Alteration in peripheral and radiculomedullary somatosensory conduction</td>
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<td></td>
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<td>(iii) Haematochemical tests</td>
<td>(iii) Motor and sensory nerve conduction abnormalities</td>
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<td>(iv) Abdominal ultrasound</td>
<td>(iv) Axonal damage</td>
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<td>(v) Monoclonal gammopathy of the IgG lambda type</td>
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<td>(vi) Gallbladder adenomyoma</td>
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<td><strong>Diagnosis</strong></td>
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<td>Cordonal syndrome and idiopathic peripheral neuropathy (unknown etiology)</td>
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<td>2007</td>
<td>(hospitalized)</td>
<td>San Raffaele Foundation Scientific Institute Hospital, Neurology, Clinical Neurophysiology and Neurorehabilitation, Milan, Italy</td>
<td>(i) Neurologicalevaluation</td>
<td>(i) Postural instability progression</td>
<td>Lack of documentation</td>
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<td>(ii) Motor and somatosensory conduction abnormalities</td>
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<td>(iii) <strong>ANA test positivity</strong>: 1: 80</td>
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<td><strong>Diagnosis</strong></td>
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<td>“Sensorymotor neuropathy of undetermined cause and spondylogenic myelopathy”</td>
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<td>paralleled by monoclonal gammopathy of undetermined significance (MGUS)</td>
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<td>2008</td>
<td>(January–March)</td>
<td>—</td>
<td>Self-evaluation</td>
<td>(i) Postural instability progression</td>
<td>Self-administration of betamethasone (2 mg/die)</td>
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<td>(ii) Motor and somatosensory conduction abnormalities</td>
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<td>(iii) <strong>ANA test positivity</strong>: 1: 80</td>
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<td><strong>Diagnosis</strong></td>
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<td>paralleled by monoclonal gammopathy of undetermined significance (MGUS)</td>
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<td>2008</td>
<td>(September, hospitalized)</td>
<td>Neurology and Neurophysiology, Poli clinico P. Giaccone Hospital, Palermo</td>
<td>(i) Motor evoked potentials (MEP) and somatosensory evoked potential (SSEP) monitoring</td>
<td>(i) Postural instability progression</td>
<td>Lack of documentation</td>
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<tr>
<td></td>
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<td>(ii) Haematochemical tests (including Antinuclear Antibody (ANA) test)</td>
<td>(ii) Motor and somatosensory conduction abnormalities</td>
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<td>(iii) <strong>ANA test positivity</strong>: 1: 80</td>
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<td>paralleled by monoclonal gammopathy of undetermined significance (MGUS)</td>
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<td>2009</td>
<td>(clinical consultation)</td>
<td>Carlo Besta Neurological Institute, Milan</td>
<td>(i) Neurological evaluation</td>
<td>Lack of documentation</td>
<td>Dexamethasone 25 mg/die</td>
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<td>2010–January 2011</td>
<td>Private consultation with a neurologist, Anversa, Belgium</td>
<td>(i) Blood mercury levels determination</td>
<td>(i) Postural instability progression</td>
<td>Chelation therapy cycles with i.v. administration of 2,3-dimercapto-1-propanesulfonic (DMPS) acid</td>
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at admission remained stable during all monitoring period, until eight months after the end of the complete chelation therapy.

Potential kidney effect was also investigated by evaluating standard renal function markers, that is, creatinine, uric acid, electrolytes, serum beta-2 microglobulin, and N-acetyl-beta-D-glucosaminidase, all resulting in healthy reference ranges, and clinical tests, for example, Giordano’s sign, resulted bilaterally negative. Other laboratory findings fell in the considered normal range (data not shown).

2.1. Methods in Brief. We periodically monitored Hg concentrations in blood and urine (BHg and UHg) from admission to our TU (t₀) until 1 year later (about 410 days) (t₂), around eight months after the end of the complete chelation therapy. For the urinary Hg determination, 24-hour urine specimens were collected.

At admission (t₀), even rare metals and trace elements (e.g., lead, zinc, magnesium, and manganese) were also measured. All the evaluated levels fell in the healthy normal ranges (data not shown).

Notably, two days after admission, a chelation challenge test with meso-2,3-dimercaptosuccinic acid (DMSA, Succicaptal®), reflecting the mobilizable and likely toxicologically active fraction of the Hg body burden, was performed (two equal oral doses every 8 hrs: 10 mg/kg/dose), strictly monitoring BHg and UHg.

2.1.1. Mercury Determination. Mercury levels were determined by inductively coupled plasma-mass spectrometry (ICP-MS) while speciation analyses were conducted using HPLC-ICP-MS [11, 12].

Accuracy was checked by reference solutions (8AB occupational G-EQUAS for blood and urine). The detection limits and the coefficients of variability (CV%) for the different matrices were 0.1 microg/L and 5% for BHg and 0.05 microg/L and 4% for UHg, respectively.

2.1.2. Blood Cells Isolation and Determination of Peripheral Neurochemical Markers, That Is, Muscarinic Receptors in Lymphocytes (l-MRs) and Monoamine Oxidase B in Platelets (p-MAO-B). For l-MR and p-MAO-B determinations in lymphocytes and in platelets, respectively, blood samples were collected into EDTA containing tube and immediately processed to isolate lymphocytes for MR binding or platelets for MAO-B activity as previously described [13]. The p-MAO-B activity was determined radiochemically in duplicate samples as described by Coccini et al. using 14C-P-PEA as the substrate [14]. Specific activity was determined in the presence of pargyline hydrochloride. The enzyme activity was expressed as nanomol/L/mg protein/h.

Muscarinic receptors were determined by binding assays using a single concentration (Kd) of the specific tritiated ligand antagonist [3H]QNB for muscarinic receptors in lymphocytes [13]. The specific binding was measured in the presence or absence of atropine. Each sample was assayed in triplicate and data were expressed as femtomol/L/10⁶ cells.

3. Results

The patient was evaluated and monitored both during the 7-day hospitalization at TU of FSM Pavia hospital and throughout the following 1-year period.

(I) t₀: at admission to our Toxicology Unit, BHg and UHg levels were 27 and 1.4 microg/L, respectively (normal values: BHg 1–4.5 microg/L; UHg 0.1–4.5 microg/L) (Figure 1). Parallelly, neurochemical markers, that is, muscarinic receptors in lymphocytes (l-MRs) and monoamine oxidase B in platelets (p-MAO-B), have been determined. The latter analyses demonstrated (i) a strong, significant increase in lymphocyte-MRs, that is, 185.82 femtomol/million lymphocytes (normal range: 8.0–16.0), and (ii) a normal p-MAO-B activity of about 10.46 nanomol/mg prot/hr (normal range: 7.0–11.0) (Figures 2(a) and 2(b)).

(II) t₀: two days after the DMSA (Succicaptal) chelation challenge test, (i) a weak reduction of BHg concentration was measured, paralleled by a 3.7-fold increase of UHg concentration. Specifically, the following BHg and UHg levels were measured: 24.5 and 5.2 microg/L, respectively (Figure 1). Elemental-Hg (before and after mobilization) and methyl-Hg (after mobilization) were evidenced by Hg speciation
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Upper normal limit value (4.5 microg/L)

![Graph](image)

**Figure 1: Hg concentration trends in different sampled matrices, that is, blood (BHg) and urine (UHg), throughout the entire biological monitoring 1-year period (from \( t_0 \) to \( t_2 \)). The hatched bars indicate the 16-h chelation challenge test and the successive two DMSA cycles.**

Accordingly, BHg and UHg concentrations were settled on normal reference value, showing negligible difference with those determined eight months before, at the end of the chelation therapy \( (t_0 + 5 \text{ months}) \) (Figure 1).

Importantly, it should be highlighted that parallel neurophysiological evaluation demonstrated a complete remission of the detrimental neurological symptoms.

### 4. Discussion

Evaluating the effects of exposure to neurotoxicants is extremely difficult in human investigations. The Hg exposure described in the present case report is quite different than that occurring in classical occupational setting. This accidental Hg vapour long-lasting exposure is hardly quantifiable, characterized by inhomogeneous, unlikely predictable indoor concentrations. If it is widely accepted that repeated and regular chronic Hg exposure (e.g., in occupational setting) causes increases of both urine and blood Hg levels, on the other hand, so far, the outcome of biomarkers expected in a peculiar environmental exposure scenario like the one described in our case report is not clearly documented. Specifically, the patient exposure to Hg in the study-room was irregular, lasting an amount of time different day by day, with Hg air concentration unpredictable and probably dissimilar in the different areas of the room. Our findings demonstrated a high BHg value already at admission, paralleled by a normal UHg level. Particularly, even though a physiological kidney function was observed (as indicated by normal renal parameters as well as clinical evaluation), unexpectedly we did not determine an enhanced Hg renal excretion at admission. We can hypothesize that Hg remained deposited in some tissues (particularly in nervous system), as demonstrated by the detrimental neurological symptoms. Moreover it is possible that this “irregular” exposure led a Hg accumulation in patient sufficiently to induce neurotoxicity without causing an evident Hg increase in urine. In this case report, Hg exposure may be considered as the consequence of repeated short exposures, some hours a day only and even not every day: this exposure pattern could justify the Hg increase in blood (as indicator of occurring exposure) associated with low Hg urine levels resulting from repeated but “irregular” and spotted exposures. In support of this hypothesis, the DMSA chelation challenge test induced a 3.7-fold increase of BHg concentration, thus demonstrating the mobilization of Hg levels from tissues and the atypical chronic accumulation mirrored by the atypical Hg urinary levels at \( t_0 \).

In this respect, the present study applying a complementary approach, which correlates specific exposure parameters (i.e., analytical data) and indicators of neural cell function, in
peripheral blood cells, supported and properly addressed a differential diagnosis, thus representing a promising strategy to be used in clinical setting.

One characteristic of the neurochemical parameters presently investigated, that is, muscarinic receptors (MRs) and monoamine oxidase activity type-B (MAO-B), is that they are also expressed in easily accessible matrices, for example, blood components, such as lymphocytes and platelets. Although novel noninvasive radiological imaging techniques such as NMR, PET, and SPECT (magnetic resonance, positron emission tomography, and single-photon emission computed tomography, resp.) may allow directly estimating MRs and MAO activity in living human brain [15], these methods are expensive in terms of cost and their widespread application to neurotoxicological investigations can not be proposed on a large scale, thus supporting the need to employ the above-mentioned peripheral neurochemical parameters, to investigate the status of homologous CNS markers [16–18].

Even though the use of biochemical markers in neurotoxicology is particularly challenging due to (i) the complexity of CNS functions, (ii) the multistage nature of neurotoxic events, and (iii) the inaccessibility of target tissue, in recent years great effort has been devoted to develop and validate new surrogate parameters in peripheral tissues easily and ethically obtained in humans, reflecting the same parameters in nerve tissue.

Specifically, peripheral blood lymphocytes are considered the main tool to explore cholinergic function, as also supported by human studies demonstrating similar immunoblotting patterns both in lymphocyte and in striatum membranes [19]. Further experimental evidence showed that MRs binding can be similarly modulated by cholinergic agonists and antagonists, in both lymphocytes and brain tissue [20].

For these intriguing peculiarities, in our previous studies, intended to validate their use as peripheral surrogate markers in experimental controlled conditions, MRs have been investigated as biomarkers of neurotoxicity in animals exposed to environmental chemicals, demonstrating to reflect analogous receptor changes occurring in rat brain after repeated MeHg exposure during adult age as well as during development [6, 21].

Additionally, peripheral MRs have been clinically applied as predictors of pharmacological response in psychotropic drugs-treated subjects, as well as to investigate the role of neurochemical disturbances in affective disorders and neurological diseases (e.g., Alzheimer’s, Parkinson’s and Meniere’s diseases, and Gilles de la Tourette syndrome), clearly demonstrating significative MRs binding alterations [16, 17, 19].

Moreover, our previous investigation showed a significant level reduction of surrogate peripheral markers of cholinergic and monoaminergic neurotransmissions in attention deficit hyperactivity disorder (ADHD) children. Particularly, a relationship has been demonstrated between l-MR binding levels and specific ADHD symptoms such as inattention and ODD in unmedicated subjects [22].

With regard to the other measured peripheral marker, that is, platelet MAO-B (p-MAO-B), it is the sole type in human platelets and the primary type in the human brain (80–95% of total MAO), playing a pivotal role in the catabolism of various neuroactive and vasoactive amines, that is, neurotransmitters (including dopamine), being located in CNS, as well as many peripheral tissues.

The amino acid sequences of MAO-B in both platelets and brain are identical and the biochemical and pharmacological characteristics of this isoenzyme are also similar in the two tissues. For these reasons, p-MAO-B activity was proposed as a predictive peripheral marker of various psychopathologies [23], neurodegenerative diseases [24], and CNS neurotoxic alterations [21]; further, alterations in MAO levels have been implicated in the pathogenesis of psychiatric disorders. As such, decreased platelet MAO-B activity was found in children with ADHD [22, 25]. Altered p-MAO-B has been also suggested as a biomarker of alcohol dependence or alcohol consumption [14, 26].

With regard to neurotoxic compounds, this platelet enzyme has also been applied as peripheral biomarker of...
monoamine neurotransmission in patients exposed to neurotoxictants such as styrene [27] or environmental Hg [28].

In summary, positive support in the use of the neurochemical markers clearly emerges from a bulk of human literature data providing typical alteration of these parameters when used (i) as peripheral indicators of the brain neurochemistry changes associated with neuropsychiatric disorders and drug dependence or (ii) as predictors of environmental neurotoxicants exposure.

Moreover, our previous papers delineate the relevant contribution of in vivo (animal and human) researches to identifying specific molecular CNS targets of neurotoxictants which can be applied as accessible tools to use in environmental medicine as well as in clinical setting for assessing and monitoring specific exposure scenarios [13, 21].

In this view, integrated investigation approach using peripheral neurochemical markers in combination with clinical neurological evaluation and analytical data (contextually considered with patient’s history) could represent a valuable methodological strategy by which human neurotoxicity assessment may become more focused, particularly in chronic exposures.

The study clearly shows an evident association among Hg exposure levels in biological specimens, blood cholinergic markers, and clinical manifestation trend. Specifically, high/low BHg and UHg levels were accompanied by high/low MRs in lymphocytes as well as severe/slight neurological symptoms.

This is the first documented case in human of a valuable application of a specific neurochemical biomarker enabling the clinicians to support specific early differential diagnosis (i.e., Hg poisoning) and to monitor the chelation therapy efficacy.

Consent

Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review.

Disclosure

The authors alone are responsible for the content and writing of the paper.

Competing Interests

The authors report no conflict of interests.

Acknowledgments

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References


