

REVIEW ARTICLE

Immunotoxicology of arc welding fume: Worker and experimental animal studies

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Abstract

Arc welding processes generate complex aerosols composed of potentially hazardous metal fumes and gases. Millions of workers worldwide are exposed to welding aerosols daily. A health effect of welding that is of concern to the occupational health community is the development of immune system dysfunction. Increased severity, frequency, and duration of upper and lower respiratory tract infections have been reported among welders. Specifically, multiple studies have observed an excess mortality from pneumonia in welders and workers exposed to metal fumes. Although several welder cohort and experimental animal studies investigating the adverse effects of welding fume exposure on immune function have been performed, the potential mechanisms responsible for these effects are limited. The objective of this report was to review both human and animal studies that have examined the effect of welding fume pulmonary exposure on local and systemic immune responses.

Keywords: Particulate matter, inhalation, welders, epidemiology, immune

A scientific panel reported increased severity, frequency, and duration of upper and lower respirators.

Arc welding is a constitute of the contract of the co tions among welders (Lockey et al., 1988). An increased mortality from pneumonia has been observed in welders and metal fume workers (Coggon et al., 1994; Hoffmaster et al., 2006; Avashia et al., 2007). Evidence suggests that ferrous and other metal fumes inhaled in the workplace may predispose workers to pulmonary infection (Palmer et al., 2003). Unfortunately, elucidation of the mechanisms that cause these observations is lacking. The objective of this manuscript was to review both human and animal studies that have examined the effect of welding fume pulmonary exposure on local and systemic immune responses. Following a summary of what is known about welding processes and the physico-chemical characteristics of commonly used welding particles, the review will delve into the health effects from occupational exposure of humans followed by a summary of what is known about immunotoxic effects/mechanisms of effect gleaned from animal model studies.

(Received 21 October 2011; revised 12 December 2011; accepted 20 December 2011)

Arc welding is a common industrial process used to join metals. Arc temperatures above 4000°C heat the base metal pieces to be joined and the consumable electrode wire or rod that is continuously fed into the weld. Most of the generated fume is derived from the consumable electrode, which is partially volatilized in the welding process (Palmer and Eaton, 2001). Vaporized metals react with air, producing metal oxide particles that are primarily of respirable size. Arc welding processes generate complex aerosols that are composed of hazardous metal fumes (e.g., manganese (Mn), hexavalent chromium (Cr6+), nickel (Ni)), and gases (e.g., carbon monoxide (CO), ozone (O_3)).

The American Welding Society has identified more than 17 metal joining processes that are used commercially (The James F. Lincoln Arc Welding Foundation, 2000). Seven processes are classified under arc welding that together account for the greatest amount of welding and filler metal deposited commercially. Each arc welding method has its own metallurgical and operational



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Abbreviations	IgM, immunoglobulin M
Mn, manganese	ROS, reactive oxygen species
Cr ⁶⁺ , hexavalent chromium	BALF, bronchoalveolar lavage fluid
	ITI, intratracheal instillation
Ni, nickel	AM, alveolar macrophage
CO, carbon monoxide	PMN, polymorphonculear leukocyte
O ₃ , ozone	Mmp12, matrix metalloproteinase-12
MMA, shielded manual metal arc welding	Cd5l, Cd 5 molecule-like
GMA, gas metal arc welding	Ccl7, chemokine (C-C motif) ligand 7
FCA, flux-cored arc welding	Cxcl5, C-X-C motif chemokine 5
MIG, metal inert gas welding	Spp1, secreted phosphoprotein 1
TIG, tungsten inert gas welding	Trem2, triggering receptor expressed on myeloid cells 2
NO, nitrogen oxide	IgG-2a, gamma-2a immunoglobulin heavy chain
CO ₂ , carbon dioxide	Igh-1a, immunoglobulin heavy chain-1a
Cr, chromium	Igh, immunoglobulin heavy locus
Fe, iron	Ccl3 and Ccl4, chemokine (C-C motif) ligands 3 and 4
MS, mild steel	T _H 1, T-helper type 1; T _H 2, T-helper type 2
SS, stainless steel	IFN-γ, interferon-γ
Al, aluminium	TNF, tumor necrosis factor
ACGIH, American Conference of Governmental Industrial	IL, interleukin
Hygienists	MCP-1, monocyte chemotactic protein-1
TLV-TWA, Threshold Limit Value-Time Weighted Average	MIP-2, macrophage inflammatory protein-2
CRP, C reactive protein	SRBC, sheep red blood cells
NK, natural killer cells	LPS, lipopolysaccharide
IgA, immunoglobulin A	Fe ₂ O ₃ , iron oxide
IgG, immunoglobulin G	re_2O_3 , mon oxide

Table 1. Common arc welding processes and descriptions.

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Process	Synonyms	Description ^a
Shielded manual metal arc welding (MMA)	Stick welding	Simplest and most widely used process, inexpensive, weld is produced by heating an arc between a flux-covered metal electrode (consumable) and the work, shielding is obtained from decomposition of the flux covering, can weld all ferrous metals in all positions.
Gas metal arc welding (GMA)	Metal inert gas welding (MIG)	High speed, economical process, weld is produced by heating with an arc between a continuous filler metal electrode (consumable) and the work, shielding comes from an externally supplied gas mixture, top quality welds produced in all metals and alloys.
Flux-cored arc welding (FCA)	_	Weld is produced by heating with an arc between a continuous filler metal electrode (consumable) and the work, shielding is obtained from a flux contained within the electrode, gas mixtures may be used for additional shielding, produces smooth sound welds of high quality.
Gas tungsten arc welding	Tungsten inert gas welding (TIG)	Weld is produced by heating with an arc between a single tungsten (non-consumable) electrode and the work, shielding is obtained from an inert gas mixture, top quality welds can be produced using this process in all metals and alloys.
Submerged arc welding	_	Heat from the submerged arc is derived from an arc between a bare metal electrode and the work, arc is shielded by a blanket of granular fusible flux which prevents emission of arc radiation, sparks, spatter, and fumes.

^aAs referenced from The James F. Lincoln Arc Welding Foundation (2000).

advantages, and may present its own potential health and safety hazard. The most common types of welding are shielded-manual metal arc (MMA), gas metal arc (GMA), flux-cored arc (FCA), gas tungsten arc, and submerged arc welding (Table 1). The composition and the rate of generation of welding fumes are characteristic of the individual welding processes, and may be affected by the welding current, shielding gases, and the technique and skill of the welder. The choice of a particular technology depends on many factors, including the thickness and type of base metal to be welded, the size and strength of the weld desired, the speed or volume of welding, the position of the material to be welded (i.e., vertical or horizontal), and cost (Martin et al., 1997). Exposure to diverse aerosols generated from different welding processes,

combined with the complexity of workplace settings, could increase the risk of exposure and the associated adverse health effects.

Welding fume characterization

Particle morphology and size

Electron microscopic analyses (Figure 1) have revealed that most particles generated during arc welding processes are arranged as chain-line agglomerates of smaller, spherical primary particles (Zimmer and Biswas, 2001; Jenkins et al., 2005; Antonini et al., 2006). Particle formation during arc welding first includes nucleation, a process by which high temperature metal vapors are transformed into primary particles, followed

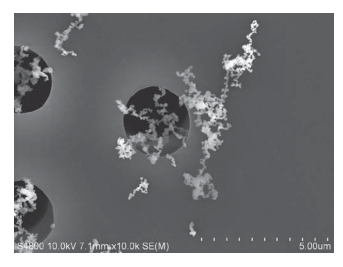


Figure 1. Scanning electron micrograph of SS welding fume collected onto a filter during welding using a flux-cored electrode. Note the chain-like agglomerates of much smaller, spherical primary particles. Welding fume samples were collected onto 47-mm Nuclepore polycarbonate filters (Whatman, Clinton, PA). The filters were cut into equal sections and mounted onto aluminum stubs with silver paste. The deposited welding particles were viewed using a JEOL 6400 scanning electron microscope (JEOL, Inc., Tokyo, Japan).

by coagulation, a dynamic aerosol growth process where smaller primary particles collide to form larger agglomerates (Zimmer, 2002). The primary particles generated during arc welding have been observed to be in the ultrafine size range, <0.10 µm (Sowards et al., 2010). Modest changes in welding process parameters (e.g., voltage) can affect the number of ultrafine welding particles formed (Antonini et al., 2011a). The agglomerates are formed after collisions between primary particleprimary particle, primary particle-agglomerate, and agglomerate-agglomerate (Zimmer, 2002). After these collisions, the agglomerates are believed held together by van der Waals, electrostatic, and magnetic forces. The mass median aerodynamic diameter of most arc welding fume is in the fine size range between 0.20-0.50 µm after reaching the welder's breathing zone.

Gas and fume chemical properties

A variety of gases are formed during the welding process that may affect the health of exposed workers (Table 2). Weld shielding gases can intensify the ultraviolet radiation produced in the arc, leading to the photochemical formation of O₂ and nitrogen oxides (NO, NO₂). Carbon dioxide (CO₂) may be reduced and converted to the chemically more stable, but potentially fatal gas, CO. The oxidation of vapors from degreasing agents (e.g., trichloroethylene), which are sometimes used to clean the base metals prior to welding, can produce the pulmonary irritant gas, phosgene.

In addition to gases, arc welding produces complex metal oxide particles that are volatilized from the consumption of the welding electrode or the flux material added within or on the outside of an electrode (Zimmer

Table 2. Potential health hazards associated with welding.

FUMES:

Alloys

Aluminum (conducive to high fume formation rates and ozone production)

Chromium (stainless steel alloy, lung carcinogen)

Copper

Iron (most common fume component, siderosis)

Lead (brass and bronze alloy, neurotoxin)

Magnesium

Manganese (neurotoxin, respiratory irritant)

Molybdenum

Nickel (stainless steel alloy, lung carcinogen)

Tin (bronze and solder alloy, metal fume fever)

Zinc (galvanized steel and paint coatings, metal fume fever)

Fluxing agents

Barium, Fluorine, Silicon, Titanium

GASES:

Ozone (O₂; lung irritant; formed when arc uv light reacts with atmospheric O₂)

Nitrogen Oxides (NO, NO₂; lung irritant; formed by oxidation of atmospheric nitrogen)

Carbon Monoxide (CO; highly toxic; at high arc temperatures CO₂ is reduced to more stable CO)

Phosgene (formed when arc uv light reacts with metal-cleaning chlorinated hydrocarbons)

RADIANT ENERGY:

Ultraviolet

Visible

Infrared

OTHER:

Heat

Noise Vibration

and Biswas, 2001). Shielding gases; base metal composition and coating; paint; the presence of degreasing, cleaning, or anti-slag chemicals; and welding process parameters may all influence the chemical composition of the resulting fume. Most welding jobs use mild or carbon and low alloy steels. Mild steel (MS) electrodes are composed predominately of iron (Fe) with low and varying amounts of Mn. Welding using stainless steel (SS), aluminum (Al), Ni, and other alloys accounts for <10% of all welding performed (Beckett, 1996). SS electrodes contain significant amounts of Cr, in addition to Fe, Mn, and Ni. Depending on the welding metals and processes used, oxides and salts of many other elements may be present in the fume (Table 2).

Based on the selection of the welding process and materials, the resulting fume may be chemically distinct. Fumes generated during GMA welding processes generally are water-insoluble and closely mimic the metal composition of the welding wire that is consumed during the process. Due to the presence of fluxes, fume formed during MMA welding has been observed to be physically and chemically more complex than fumes generated during GMA welding (Zimmer and Biswas, 2001; Jenkins and Eagar, 2005; Jenkins et al., 2005). Because of the presence of alkali metals (e.g., potassium and sodium) in the fluxes, the fumes from MMA welding processes are very water-soluble (Antonini et al., 1999).



Recent studies indicate that a metal oxidation state is highly dependent on welding process and shield gas composition (Keane et al., 2009, 2010). Mn species, including Mn2+, Mn3+, and Mn4+, have been observed in the generated fume from multiple processes, and the proportions of the different Mn species were dependent on the welding process and settings (Minni et al., 1984; Keane et al., 2010). Similarly, the concentration of Cr⁶⁺ can vary significantly depending on the welding process and shield gas type (Keane et al., 2009). These differences in metal solubility and valence states in fumes generated from different welding processes have been shown to influence the bioavailability of the metals after deposition in the respiratory tract and the eventual translocation of the metals to the circulation and other organ systems in animal models.

Using neutron-activation, Lam et al. (1979) demonstrated that the metal constituents of SS welding fumes generated from MMA and GMA processes were cleared in different phases. In the initial phase, intact welding particles cleared from the body within days up to a week by mucociliary and lung macrophage clearance. The clearance rates of each element of the SS fume were similar during this initial phase, indicating that the eliminated particles were transported in their entirety, without separation of the constituents. At the later phase, various elements of the specific fumes cleared at much different rates with biological half-lives of several weeks, indicative of tissue solubilization of the deposited particle. Previous studies by our group indicate that SS welding fumes cleared from the lungs of rats at a slower rate compared to MS fumes (Antonini et al., 1996, 2011b). In addition, specific metals in the different welding fumes cleared from the lungs at varying rates after exposure. The more toxic metals (e.g., Mn, Cr) present in some welding fumes were found to clear more quickly and completely from the lungs compared to Fe, likely increasing their translocation from the respiratory system to other organs (Antonini et al., 2010).

Welding exposure and health effects

An estimated 430,000 workers are employed full-time in welding operations in the United States (U.S. Bureau of Labor Statistics, 2010), and millions more worldwide are exposed to welding aerosols daily. Welders are part of an heterogeneous workforce employed in a variety of workplace settings; these may include well-ventilated indoor and open-air sites or confined poorly ventilated spaces (e.g., ship hulls, pipelines, building crawl spaces). Because of this, a wide range of exposure concentrations have been measured in workplaces where welding occurs. Personal exposure measurements of ≈200 workers from numerous occupational settings throughout the US indicated that total welding particulate levels ranged from 1.02–37.30 mg/m³ in 1995 and 0.10–18.00 mg/m³ in 1996 (Susi et al., 2000). In the assessment of a construction site, Meeker et al. (2007) observed total welding particulate

concentrations to range from 2.65–11.6 mg/m³ in settings without local exhaust ventilation, and 3.15-5.44 mg/m³ where local exhaust ventilation was used. Airborne fume concentration levels of individual metals in eight welding companies that mostly performed GMA-MS welding were measured by the Workplace Safety and Health Branch in Manitoba, Canada (Korczynski, 2000). The personal exposures to Fe and Mn ranged from 0.04-16.29 mg/m³ and 0.01-4.93 mg/m³, respectively. Studies have reported that workplace levels of individual metals in welding fumes have exceeded airborne exposure limits. Korczynski observed that 19% of welders studied were exposed to levels of Fe that exceeded the American Conference of Governmental Industrial Hygienists (ACGIH) Threshold Limit Value-Time Weighted Average (TLV-TWA) of 5 mg/m³. Moreover, 62% of the welders studied were reportedly exposed to levels of Mn that exceeded the ACGIH TLV-TWA of 0.2 mg/m³. In the assessment of nine construction sites among different welding trades, it was observed that the probability of exceeding the ACGIH TLV-TWA for Mn was sufficiently high to cause health concerns among the construction trades (Rappaport et al., 1999).

Hundreds of studies have evaluated the health effects associated with welding fume inhalation. However, these effects are sometimes difficult to assess because of differences in worker populations, work area ventilation, welding processes and materials used, and other occupational exposures besides welding fumes. It has been reported that most career welders experience some type of respiratory disorder during their time of employment (Sferlazza and Beckett, 1991; Martin et al., 1997; Antonini, 2003). Pulmonary effects have included metal fume fever, bronchitis, siderosis, reversible lung function decrements, increased susceptibility to infection, and an increased risk of lung cancer. In addition, a variety of non-pulmonary effects (e.g., neurological, cardio-vascular, and reproductive) have been observed in workers exposed to welding fumes. The occupationalrelated health effects of greatest concern are the potential development of neurological effects, lung cancer, and decrements in immune function and defense.

Welding fume exposure may be associated with the development of neurological dysfunction similar to Parkinson's disease. Much of this concern has been attributed to Mn in the generated welding fume. Mn intoxication following chronic human exposure in other occupational settings, such as mining, smelting, ferroalloy, and dry battery industries, has been documented to cause a Parkinson's disease-like syndrome called 'manganism' (Roels et al., 1992; Mergler and Baldwin, 1997, Lucchini et al., 1999). Neurobehavioral changes have been reported in exposed welders (Bowler et al., 2007a, b; Ellingsen et al., 2008), and case reports show Mn accumulates in dopaminergic brain regions of welders exposed to high concentrations of welding fume (Sadek et al., 2003; Josephs et al., 2005). In addition, some studies describe a potential link between welding and Parkinsonism, and

suggest the possibility of an early-onset Parkinsonism among welders (Racette et al., 2001, 2005), whereas other studies do not (Goldman et al., 2005; Stampfer, 2009; Tanner et al., 2009). Recent animal studies demonstrated that repeated exposure to Mn-containing welding fumes modulated molecular factors associated with synaptic transmission, oxidative stress, neuro-inflammation, and mitochondrial dysfunction in dopaminergic targets in rat brains, as well as induced transcriptional changes in a variety of PARK genes (Sriram et al., 2010a, b).

Despite inadequate evidence in humans and laboratory animals, welding fume has been classified as 'possibly carcinogenic' by the International Agency for Research on Cancer due to the presence of Cr⁶⁺ and Ni in SS fumes (IARC, 1990). Multiple epidemiology studies have reported an elevated risk for the development of lung cancer among welders (Hansen et al., 1996; Moulin, 1997; Becker, 1999), whereas others have not (Steenland et al., 1991; Hansen et al., 1996; Danielsen et al., 2000). Moreover, epidemiological studies have been unable to correlate the development of lung cancer solely with exposure to Cr-containing SS welding fume as compared to non-Cr-containing MS fume (Moulin et al., 1993; Langard, 1994; Moulin, 1997). However, in vitro genotoxicity studies demonstrated that Cr-containing SS, but not MS, welding fumes are mutagenic and induce DNA damage (Hedenstedt et al., 1977; Maxild et al., 1978). Recent animal studies with MMA- and GMA-generated fumes indicate these to be weak lung tumor initiators in A/J mice, a lung tumor susceptible mouse strain. However, a chronic lung inflammatory response to these fumes was found, which may suggest that long-term exposure could increase lung tumor incidence and/or multiplicity in the A/J model (Solano-Lopez et al., 2006; Zeidler-Erdely et al., 2008, 2011a, b).

Human immunotoxicity studies

Data from the national analyses of occupational mortality for England and Wales have consistently demonstrated that welders have an increased mortality from pneumonia (Registrar General 1958, 1971, 1981). In 1994, Coggon et al. (1994) confirmed these observations using 1979-1980 and 1982-1990 data from the UK that showed an excess of pneumococcal and unspecified lobar pneumonia in men below retirement age of 65 years. Lobar pneumonia, a disease with acute progression, is usually the result of infection by Streptococcus pneumoniae and affects a large and continuous area of a lung lobe. Since a similar mortality pattern was found in other metal fume-exposed workers such as molders and coremakers, Coggon et al. concluded that the primary causative agent could be the metallic fume components or possibly O₃ or NO. Studies from the US associating increased mortality from pneumonia due to welding fume exposure are few, but most support an increased risk in welders of pneumonias other than bronchopneumonia (Puntoni et al., 1979; Beaumont and Weiss, 1980; Polednak, 1981; McMillan

and Pethybridge, 1983; Milham, 1983; Newhouse et al., 1985).

In 2003, a hospital-based case-control study in males (mean age of 46 years) by Palmer et al. (2003) determined that inhalation of metal fumes renders the worker susceptible to infectious pneumonia, primarily of the lobar type, and that this was a reversible phenomenon after exposure ceased. Further, the study identified pneumonia of pneumococcal as well as Legionella, Mycoplasma, and *Haemophilus influenza* origin in the subjects. Also, ferrous metal (with or without other metals/alloys) was found to be associated with a higher risk of pneumonia than non-ferrous exposures. Importantly, the study eliminated other workplace hazards such as coal dust, wood dust, cement dust, and asbestos as contributing to the increased risk in the metal dust-exposed subjects (i.e., welders, molders, coremakers, and die casters) (Palmer et al., 2003). These results led Palmer et al. to postulate that ferrous metal fumes may cause oxidative damage to lung host defense or that they may provide an enriched free Fe environment for micro-organisms to temporarily prosper such as that which occurs with sickle cell disease (Barrett-Connor, 1971). Moreover, Ghio and Cohen (2005) have theorized that the immunomodulatory effect of particulate matter, such as welding fume, may involve a disruption of Fe homeostasis in cells and tissues. It is possible that other metals may compete with Fe so that lung phagocytes become Fe-insufficient and unable to function properly, while simultaneously allowing for greater free Fe to remain available for pathogens (Cohen et al., 2010). A subsequent study by Palmer et al. (2006) revealed that, surprisingly, despite high Fe loads and a low unsaturated Fe-binding capacity in sputum, no overt inflammatory response was found in blood or sputum of welders with a median employment of 10 years. However, an increase in blood eosinophil and basophils in welders vs non-welders, with some correlation to exposure level, was observed. Also found were trends for increased C reactive protein (CRP), neutrophil oxidative burst, sputum IgA, and decreased sputum eosinophils (Palmer et al., 2006).

Supportive evidence for metal particulate-induced epithelial damage or macrophage activation causing weak systemic effects via cytokine signaling, however, could not be found in the airways of the patients. Taken together, the findings of this study were difficult for the authors to resolve given the strong evidence for pro-inflammatory lung effects of a variety of metal-rich particulates such as diesel exhaust, residual oil fly ash, particulate matter, zinc oxide, and coal dust (Blanc et al., 1991; Carter et al., 1997; Salvi et al., 2000; Vallyathan et al., 2000; Ulrich et al., 2002). The authors speculated that the high transitional metal levels in the lung must somehow impair lung host defense and provide a temporary window for infection. Alternatively, apoptosis, via Fe-catalyzed oxidative stress, may impair immune cell activation, which may also lead to increased worker susceptibility (Palmer et al., 2006). In fact, apoptosis is a



mechanism by which *S. pneumonia*, the organism presumably at most infective risk to metal fume workers, acts to augment its own pathogenicity (Hakansson et al., 1996).

Case studies of rapidly progressing, fatal pneumonia have been reported in four healthy male welders residing in Texas and Louisiana (Miller et al., 1997; Avashia et al., 2007). All four cases exhibited symptoms of Bacillus anthracis respiratory disease, and the welders succumbed to the illness 5-8 days after their initial onset of symptoms. The causative agent was determined to be Bacillus cereus, an organism occasionally associated with food-borne illness and merely considered a contaminant in clinical laboratory specimens. In healthy individuals, B. cereus poses no health concern, but could cause a variety of infections in those who are immunocompromised or possess an underlying illness (Drobniewski, 1993). It was later discovered that three of the four B. cereus isolates obtained from the patients and/or workplace contained B. anthracis toxin genes, an unusual and unexpected finding (Miller et al., 1997; Hoffmaster et al., 2004; Avashia et al., 2007). In all cases, including another similar non-fatal case involving the same organism, the men were only linked by their metal-working profession (Hoffmaster et al., 2004). This association further raises the possibility that exposure to metal fumes may lead to periods of increased susceptibility, even to ubiquitous organisms such as *B. cereus*.

Immunological screening studies in welders are limited and, in general, report moderate changes for various immune-related parameters. In a cohort of 11 welders and nine grinders (mean age = 31 years; work experience = 8 years), Borska et al. (2003) reported decreased levels of IgG, IgA, and IgM as well as acute phase proteins transferrin, α_1 -antitrypsin, β_2 -macroglobulin, haptoglobulin, and ceruloplasmin in the serum. Increased levels of phagocytic cells, C3 complement component, neopterin, β_{\circ} -macroglobulin, and orosomucoid were also found. Others have reported similar findings of alterations in certain parameters of humoral immunity including decreased IgG, IgA, and increased C3 complement component (Boshnakova et al., 1989, Nielsen et al., 1993; Hanovcova et al., 1998). In contrast, Tuschl et al. (1997) observed no effect on lymphocytic sub-populations, mixed lymphocyte reactions, leukocytes phagocytosis, or immunoglobulin levels in an examination of 30 welders who worked in a plant where levels of welding fumes typically exceeded Austrian Working Limits of 5 mg/m³. However, the authors reported that welding fumes reduced the cytotoxic activity of lymphokine-activated killer cells which are primarily natural killer (NK) cells. They suggested that the decreased immunocompetent state might subject the welders to an increased risk of infection because NK cells are a main defense mechanism against viruses (Tuschl et al., 1997). In a cohort of 74 healthy welders, Boshnakova et al. (1989) found evidence for impaired T- and B-lymphocytes and decreased serum IgG and total E-rosette-forming cells. The authors suggested a link to complex occupational factors, such as the heavy metal Mn, noise, and vibration, as possible causes for the immune suppression.

Other studies have reported effects on lymphocytes in welders. For example, a study examining the effects of Mn exposure in welders (work experience = 6-36 years) found a significant inverse correlation between blood Mn concentrations and CD8+, CD3+, CD4+, naïve T, and total lymphocytes after adjusting for age and smoking (Nakata et al., 2006). In addition, B (CD19⁺)-lymphocyte numbers were also significantly decreased in high Mn-exposed welders. In a 3-year study by Hanovcova et al. (1998), increased lymphocyte counts, T-lymphocyte percentages, leukocyte metabolism, and decreased phagocytosis were found in SS welders (n = 22-53) compared to a standard laboratory reference value/non-exposed persons from the same plant. Exposure to metal fumes has been shown to induce systemic inflammation and oxidative stress parameters in the serum of welders, which could potentially lead to immune effects (Comhair and Erzurum, 2002; Han et al., 2005; Kim et al., 2005; Fang et al., 2009). A recent study by du Plessis et al. (2010) measured the oxidative status of peripheral blood mononuclear cells in welders (n = 15, mean age = 32 years, work experience = 0.25-29 years) from a South African manufacturing factory using a GMA welding process. Using flow cytometry, increased levels of reactive oxygen species (ROS) and lipid peroxidation, along with a decrease in intracellular glutathione were found. These data further support a role for an altered oxidative status in the immune suppression that occurs in workers exposed to metal fumes.

Animal immunotoxicity studies

Numerous animal studies have examined the toxic effects of welding fumes generated from MMA and GMA welding using different pulmonary exposure routes, such as pharyngeal aspiration, intratracheal instillation (ITI), and inhalation (Taylor et al., 2003; Sung et al., 2004; Yu et al., 2004; Antonini et al., 2007). Overall, analysis of the bronchoalveolar lavage fluid (BALF) has shown significant cytotoxicity, air-blood barrier damage, and an inflammatory cytokine response after welding fume exposure via these routes. In addition, a cellular influx consisting of alveolar macrophages (AM), neutrophils, lymphocytes, and, to a lesser extent, eosinophils was reported. Studies in both rats and mice that directly compared different types of welding fumes showed that SS fumes induced more lung toxicity as compared to MS fumes, an effect likely attributable to the greater proportion of toxic metals, such as Cr (Taylor et al., 2003; Zeidler-Erdely et al., 2008; Erdely et al., 2011b).

Results of lung gene expression studies in both mice and rats indicate a strong immunologic transcriptional response to welding fume (Oh et al., 2009, 2011; Yang et al., 2009; Zeidler-Erdely et al., 2010). A 30-day inhalation exposure to MMA-SS fume in rats resulted in marked changes in several immune-related genes, including



matrix metalloproteinase-12 (Mmp12), Cd 5 moleculelike (Cd5l), chemokine (C-C motif) ligand 7 (Ccl7), C-X-C motif chemokine 5 (Cxcl5), and secreted phosphoprotein 1 (Spp1). In addition, triggering receptor expressed on myeloid cells 2 (*Trem2*), γ-2a immunoglobulin heavy chain (IgG-2a), immunoglobulin heavy chain-1a (Igh-1a), and immunoglobulin heavy locus (Igh) were increased after an exposure and recovery protocol. A second 30-day MMA-SS fume inhalation exposure verified that this gene sub-set was predominant in the response and that the lung showed a lack of adaptation to a repeated exposure (Yang et al., 2009; Oh et al., 2011). In mice, many of these same genes were altered after GMA welding fume aspiration and inhalation exposure (Zeidler-Erdely et al., 2010, 2011c), which suggests a consistency across rodent species and a general response to welding fume-induced lung injury. While only portions of a GMA-SS fume inhalation mouse study (40 mg/m³ for 10 days at 3 h/day with 110-120 µg measured total deposition) have been published (Erdely et al., 2011a; Zeidler-Erdely et al., 2011c), the preliminary analysis indicates a robust immunologic response through 28 days post-exposure. The functional analysis of pulmonary gene expression changes indicated 'immune cell trafficking', 'infectious disease', 'immunological disease, and 'anti-microbial response' as highly significant categories of response. In addition, significant canonical pathways including 'communication between innate and adaptive immune cells, 'dendritic cell maturation, 'role of pattern recognition receptor in recognition of bacteria and viruses, and 'crosstalk between dendritic cells and NK cells' suggest prominent immune-mediated signaling. Further, aspirated GMA-SS welding fume (85 μg × 4 doses) in mice also increased lung gene expression of chemotactic and immunomodulatory genes such as chemokine (C-C motif) ligands 3 and 4 (Ccl3 and Ccl4), immunoglobulin M (IgM), and Mmp12 through 16 weeks post-exposure. Overall, the study concluded that sustained lung inflammation was associated with alterations in immunomodulatory genes, indicating that a delayed resolution of inflammation was linked to ongoing immune signaling in mice (Zeidler-Erdely et al., 2010).

Several animal studies have examined the immunosuppressive effects of welding fume and specifically analyzed the ability of the rat lung to clear a bacterial infection to *Listeria monocytogenes* after welding fume exposure. A preliminary study compared three fumes, GMA-SS, GMA-MS, and MMA-SS, administered by ITI (2.00 mg/rat) 3 days before pulmonary inoculation, and the subsequent effects on the lung host defense response in rats (Antonini et al., 2004). In this study, only the MMA-SS fume resulted in significant weight loss and a reduction in pulmonary clearance of L. monocytogenes after infection. However, bacterial challenge 3 days after welding fume treatment also resulted in enhanced pulmonary polymorphonuclear leukocytes (PMN) influx, ROS production, and BALF NO production. Interestingly, these factors have been suggested to be mechanisms by which bacterial clearance was enhanced after silica exposure (Antonini et al., 2000), but findings with MMA-SS fume suggest only a minor role for these parameters in the effective clearance of a bacterial challenge.

Pre-exposure to MMA-SS fume (2.00 mg/rat) before pulmonary L. monocytogenes challenge has also been shown to alter the induction of several immune-related cytokines. For instance, pre-treatment with MMA-SS fume prevented any L. monocytogenes-induced increases in interleukin (IL)-2 and also augmented IL-10 production after infection (Antonini et al., 2004). IL-10 is an antiinflammatory cytokine produced by various cell types of the innate and adaptive immune response. IL-10 reduces tissue injury of exacerbated immune responses and the expression of pro-inflammatory cytokines (Ouyang et al., 2011). While protective, the increased IL-10 may also be a mechanism of immune evasion of bacterial and viral infections (Ouyang et al., 2011). IL-2 induces responses that are both inflammatory-generating T-helper type 1 (T_H1) and T_H2 effector cells, and anti-inflammatoryincreasing the survival and functionality of regulatory T-lymphocytes and inhibiting of T₁17 differentiation (Hoyer et al., 2008). Interestingly, IL-2-deficient mice have increased T_H1 responsiveness and develop autoimmunity and inflammation, a consequence that may be linked to reduced regulatory T-lymphocytes (Hoyer et al., 2008). These studies indicate that IL-2 plays a homeostatic role in an inflammatory immune response. The reduced IL-2 response with MMA-SS fume treatment alone and before L. monocytogenes challenge suggests altered T-cell activity.

The majority of the inflammatory effects produced by MMA-SS welding fume can be attributed to the soluble, Cr-abundant component (Taylor et al., 2003; Antonini and Roberts, 2007). The immunosuppressive effects of Cr are well documented (Cohen, 2004). For example, a study that examined pre-treatment with the total MMA-SS fume and individual metal components before infection showed that soluble Cr in the fume, as opposed to the Ni or Fe, resulted in reduced bacterial clearance and excessive lung injury in rats (Antonini and Roberts, 2007). In contrast to the total fume, there was no effect on IL-10 with soluble Cr alone, which suggests IL-10 may not play a major role. IL-2 levels were reduced by soluble Cr treatment alone and pre-treatment before bacterial infection, suggesting an altered T-lymphocyte activation; this result is consistent with MMA-SS fume exposure (Antonini et al., 2004; Antonini and Roberts, 2007). Pre-exposure with soluble Cr greatly enhanced infection-induced IL-12 production in the BALF (Antonini and Roberts, 2007). IL-12 aids in the development of cell mediated immunity by increasing interferon-γ (IFNγ) production, differentiating T-lymphocytes into T_H1 effector cells, and enhancing the cytotoxicity of NK and cytotoxic T-lymphocytes (Gee et al., 2009). While this result suggested an enhanced ability for bacterial lung clearance, this was not the case in the study by Antonini and Roberts (2007).

The development of the robotic welder at the National Institute for Occupational Safety and Health (Antonini et al., 2006) has allowed the effects of inhalation of GMA-SS



and GMA-MS welding fumes with L. monocytogenes challenge to be studied in rats. Unlike in the studies that used ITI, direct inhalation (15 mg or 40 mg/m³ \times 3 h/day \times 10 days) of both of these welding fumes resulted in reduced body weight and bacterial clearance after infection (Antonini et al., 2007, 2009). Following GMA-SS fume exposure, the reduced bacterial clearance was accompanied by marked enhancement of tumor necrosis factor (TNF)-α, IL-6, IL-10, IL-12, monocyte chemotactic protein-1 (MCP-1), and macrophage inflammatory protein-2 (MIP-2) and a significant reduction in IL-2 after infection compared to air controls (Antonini et al., 2007). The reduction in IL-2 and increased IL-12 was consistent with MMA-SS fume and/or soluble Cr. Interestingly, and in contrast to MMA-SS fume and soluble Cr, GMA-SS fume inhalation reduced ROS production from AM measured via chemiluminesence. Attenuated AM activity has also been found in conjunction with reduced bacterial clearance after diesel and O₂ exposure (van Loveren et al., 1988; Cohen et al., 2002; Yin et al., 2002). Inhalation exposure to GMA-MS fume (40 mg/m 3 × 3 h/day × 10 days) did not result in an overt lung inflammatory response, but significantly reduced pulmonary bacterial clearance. In addition, pre-exposure to GMA-MS welding fume only enhanced TNF α and MIP-2 following infection (Antonini et al., 2009). Although differences were not significant, IL-2 and IL-12 responses were similar compared to SS welding exposures.

Findings from studies with *L. monocytogenes* challenge were inconsistent. The differences between aspiration and ITI of GMA fumes may be the result of AM function and cellular influx. After ITI exposure, there was an immediate and abundant influx of PMN that was delayed or absent with inhalation (Taylor et al., 2003; Antonini et al., 2007, 2009). The delayed response after inhalation to GMA fumes was attributed to inhibitory effects on AM chemotactic signaling (Antonini et al., 2007, 2009). The lack of an initial PMN influx could diminish the effectiveness of the lung to clear bacteria post-exposure. The question then becomes, why does MMA-SS fume exposure, which resulted in a significant early PMN influx, result in reduced bacterial clearance? Perhaps the answer lies between the effectiveness of local and extravasating leukocytes and the altered pulmonary cytokine response after L. monocytogenes challenge. As discussed later, circulating leukocytes exhibit a diminished response to a secondary stimulus after MMA-SS fume exposure (unpublished observations). Cohen et al. (1998) also showed that AM harvested from rats exposed to Cr had a reduced response ex vivo when stimulated with lipopolysaccharide (LPS).

These studies indicate reduced effectiveness of local and circulating leukocytes after exposure.

GMA-SS fume-exposed rats had a greater post-infection inflammatory response (Table 3), larger reductions in body weight, and poorer bacterial clearance (Figure 2) than GMA-MS fume-exposed rats. While only qualitatively comparable because of different exposure routes, the fold increase of bacteria colony forming units due to MMA-SS fume exposure was 1.6-1.8 (Antonini et al., 2004; Antonini and Roberts, 2007). Therefore, there is a graded response in reduced immune function for the different types of welding fume that corresponds to the level of general cytotoxicity and

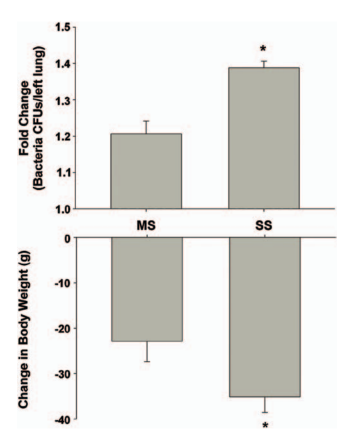


Figure 2. Comparison of the response to L. monocytogenes following gas metal arc (GMA)-mild steel (MS) or -stainless steel (SS) welding fume pre-exposure. Rats were exposed by inhalation to either GMA-MS or GMA-SS fume at 40 mg/m3 for 3 h/day for 3 days and then challenged with L. monocytogenes infection. At 3 days post-infection, colony-forming units (CFU) increased significantly and body weight decreased with exposure to either fume (adapted from Antonini et al., 2007, 2009). When represented as fold change from respective air-exposed L. monocytogenes challenged sham, there was a greater effect following GMA-SS welding fume exposure compared to GMA-MS; *p < 0.05 MS vs SS.

Table 3. Enhanced cytokine response to GMA-SS welding fume compared to GMA-MS.

	IL-2	IL-6	IL-12p70	MCP-1	MIP-2
GMA-MS	0.88 ± 0.15	0.54 ± 0.27	1.60 ± 0.27	1.15 ± 0.10	1.53 ± 0.18
GMA-SS	$\boldsymbol{0.49 \pm 0.04}$	6.05 ± 0.44	1.72 ± 0.40	4.42 ± 0.37	2.84 ± 0.47
<i>p</i> -value	0.046	< 0.001	0.803	0.001	0.071

Fold change in lavage fluid cytokine levels 3 days after L. monocytogenes infection in rats pre-exposed to 40 mg/m3 of gas metal arc welding (GMA)-mild steel (MS) or GMA-stainless steel (SS) for 3 h/day for 3 days compared to respective infection challenged air-exposed shams. Comparisons were made from data adapted from Antonini et al. (2007, 2009).



inflammation (SS > MS). The specific mechanisms responsible for reduced immune competence after GMA-MS fume exposure are unknown. Possibly, Fe, the primary metal component of the fume, promotes general bacteria viability. While plausible, exposure to iron oxide (Fe₂O₂) had no effect on bacterial clearance in rats (Antonini and Roberts, 2007). In comparison, both SS fumes resulted in a further reduced capacity to manage a bacterial challenge compared to GMA-MS fumes. These effects are likely linked to the differences in metal composition between the fumes. Cr in the SS fumes is the most suspect because of its immunosuppressive actions (Cohen, 2004). Bacterial challenge after SS exposure in rats resulted in alterations of several cytokines linked to innate and adaptive immunity not significantly altered with MS exposure. In addition, the inability of these SS-exposed hosts to efficiently respond to and clear L. monocytogenes (as compared to infected shams) caused an augmented pulmonary toxicity, likely exacerbating the situation.

In addition to the localized effects, a few animal studies provide evidence of systemic immunosuppression. One study evaluated the local and systemic humoral immune response following aspiration of multiple doses (5-20 mg/kg body weight every 5 days \times 4 doses) of MMA-SS fume in mice (Anderson et al., 2007). In vitro exposure of splenocytes to MMA-SS fume reduced IgM activity in response to sheep red blood cells (SRBC), which was the result of the soluble fume fraction. After an in vivo exposure to this fume, splenocytes also showed a significant decrease in the IgM response to SRBC, but only at the highest dose of 20 mg/kg. Localized immunosuppression was determined in B-lymphocytes harvested from lung-associated lymph nodes. There was a dose response of decreased IgM for the total and soluble fraction of MMA-SS fume, but not the insoluble fraction. The total number of B- and T-lymphocytes in the lungassociated lymph nodes increased, indicating the effect was the result of reduced function.

Recently, Antonini et al. (submitted companion paper) examined the effect of ITI pulmonary treatment of two doses of welding fumes that contained different Mn levels on the systemic immune cell profile in rats. Systemic PMN were increased, in most cases, after single or repeated ITI exposure to either a high (2.00 mg/rat) or a low (0.125 mg/rat) Mn welding fume. In terms of circulating lymphocyte number, the high Mn welding fume caused the greatest effect on the systemic immune T-lymphocyte differential, whereas neither dose of the low Mn fume significantly changed the number of circulating total T-lymphocyte or T-lymphocyte sub-populations. Significant reductions in circulating total T-lymphocytes and sub-sets were observed after ITI treatment with a 2.00 mg/rat dose of the high Mn fume. The response was similar, regardless of whether the animals were exposed to 2.00 mg/rat of the high Mn fume via single or repeated ITI treatments, indicating that the effect was acute and persisted over time. Changes in the number of whole blood B-lymphocytes or in circulating levels of IgG,

IgA, and IgM were not found after treatment with either welding fume. Importantly, a high repeated pulmonary dose (2.00 mg/rat ITI, once a week for 7 weeks) of pure manganese chloride (MnCl $_2$) did not cause a significant decrease in circulating T-lymphocytes and lymphocyte sub-sets. This indicated that the immunomodulatory effects from the parent welding fumes were likely due to pulmonary exposure to a combination of several fume-associated metals, and not solely dependent on Mn.

Unpublished experiments from our group have revealed a reduced capacity of circulating leukocytes to respond to a second stimulus after welding fume exposure. Rats were exposed to MMA-SS and sacrificed 24 h post-exposure. Whole blood was collected and incubated in sterile media with or without LPS. After 24-h incubation, supernatants were collected for cytokine analysis. Despite the increased total number of circulating leukocytes at 24 h post-exposure, LPS-induced inflammatory proteins were significantly less in MMA-SS fume-exposed rats (unpublished observations). Ongoing studies are analyzing whole blood cell gene expression by microarray after exposure and *ex vivo* stimulation to assess mechanisms related to this depressed activity.

Rim et al. (2004) used suppression-subtractive hybridization and a cDNA microarray to determine expression changes in rat mononuclear cells after 30 days of host exposure to MMA-SS fume by inhalation (108 mg/m 3 × 2 h/day × 30 days). The overall finding of this study was that MMA-SS fume down-regulated ~3-times more genes than were increased. The result was suggested to be in response to the toxicity of the prolonged exposure to MMA-SS fume. Together, these studies addressing systemic immunosuppression strongly suggest a reduced capacity to fight off an infection after welding fume exposure, in particular MMA-SS fume, both locally and systemically.

A recent study showed that mice exposed by inhalation of GMA-SS fume ($40~\text{mg/m}^3$ for 3 h/day for 10 days—daily deposition of 11.8 µg/day) resulted in local and systemic immune-related responses (Erdely et al., 2011a). This study analyzed whole blood cells, aorta, and lung by microarray, and used pathway analysis to identify consistent mechanisms of effect across tissue types. Interestingly, the major functional categories in the vasculature and blood cells were related to infectious mechanism/disease (Table 4). Systemically, the response

Table 4. Functional analysis of extra-pulmonary effects following GMA-SS inhalation.

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Aorta	Whole blood cell
Antimicrobial Response	Antimicrobial Response
Inflammatory Response	Inflammatory Response
Infection Mechanism	Immunological Disease
Immunological Disease	Infectious Disease
Inflammatory Disease	Infection Mechanism

The top five functional categories in the aorta and whole blood cells following Ingenuity Pathway Analysis of global transcriptional changes in mice exposed to gas metal arc welding-stainless steel (GMA-SS) by inhalation at 40 mg/m 3 for 3 h/day for 10 days. Data was adapted from Erdely et al. (2011a).



to GMA-SS fume appeared to be as if the particle was an infection, a response that was initiated by the lung. The major finding of the study was a prominent Type I IFN response mediated by IRF7-dependent signaling that was consistent across tissue and time. Type I IFN receptor knockout mice are resistant to the effects of L. monocytogenes (Auerbuch et al., 2004), which may provide an additional mechanism by which GMA-SS fume elicited enhanced adverse response to infection compared to GMA-MS fume (Figure 2, Table 3).

Summary

The majority of epidemiological studies suggest a reduced ability of workers exposed to metal fumes to ward off a pulmonary infection, thus allowing a window of opportunity for pulmonary infection shortly after or during a metal fume exposure (Table 5). These studies are well supported by infectivity studies in animal models (Table 6). Mechanistic studies suggest a diminished functionality of local and circulating immune cells. The reduced

Table 5. Human studies related to welding fume exposure and immune function.

Table 5. Human studies related to weld	ng tume exposure and immune function.	
Author	Finding	
Avashia et al. (2007)	Fatal pneumonia case in male metalworkers in Texas	
Beaumont and Weiss (1980)	Excess mortality from pneumonia with welding fume exposure	
Borska et al. (2003)	Alterations of systemic immunologic proteins	
Boshnakova et al. (1989)	Suppression of T- and B-lymphocyte immune system	
Coggon et al. (1994)	Pneumococcal and unspecified lobar pneumonia in metal workers	
Comhair and Erzurum (2002)	Welding fume exposure associated with systemic inflammation and/or oxidative stress	
du Plessis et al. (2010)	Welding fume exposure associated with systemic inflammation and/or oxidative stress	
Fang et al. (2009)	Welding fume exposure associated with systemic inflammation and/or oxidative stress	
Registrar General (1958, 1971, 1981)	Welders have increased mortality from pneumonia	
Han et al. (2005)	Welding fume exposure associated with systemic inflammation and/or oxidative stress	
Hanovcova et al. (1998)	Alterations of systemic immunologic proteins	
Hoffmaster et al. (2004)	Non-fatal case of pneumonia in a male metalworker in Louisiana	
Kim et al. (2005)	Welding fume exposure associated with acute systemic inflammation and/or oxidative stress	
McMillan and Pethybridge (1983)	Mortality from pneumonia with welding fume exposure	
Milham (1983)	Mortality from pneumonia with welding fume exposure	
Miller et al. (1997)	Fatal pneumonia case of two male metalworkers in Louisiana	
Nakata et al. (2006)	Inverse relationship between blood manganese levels and lymphocyte populations	
Newhouse et al. (1985)	Mortality from pneumonia with welding fume exposure	
Nielsen et al. (1993)	Alterations of systemic immunologic proteins	
Palmer et al. (2003)	Inhalation of metal fumes renders the worker susceptible to infectious pneumonia	
Palmer et al. (2006)	Chronic metal fume exposure impairs lung defenses	
Polednak (1981)	Mortality from pneumonia with welding fume exposure	
Puntoni et al. (1979)	Mortality from pneumonia with welding fume exposure	
Tuschl et al. (1997)	Welding fume reduced the cytotoxic activity of lymphokine-activated killer cells	

Table 6. Animal studies related to welding fume exposure and immune function

Author	Welding fume	Animal model	Finding
Route of Exposu	re: Inhalation		
Antonini et al. (2007)	GMA-SS; 15 or 40 mg/m 3 × 3 h/day × 1, 3, or 10 days; inoculation 1 day later with 5 × 10 4 <i>L. monocytogenes</i>	Sprague-Dawley male rats	GMA-SS welding fume induced lung injury, reduced bacterial clearance with a delayed effect on pulmonary inflammation
Antonini et al. (2009)	GMA-MS; $40 \text{ mg/m}^3 \times 3 \text{ h/day} \times 3 \text{ or } 10$ days; inoculation 1 day later with $5 \times 10^4 L$. monocytogenes	Sprague-Dawley male rats	GMA-MS had no effect on parameters of lung injury and inflammation but reduced bacterial clearance
Antonini et al. (2011b)	GMA-MS, GMA-SS; $40 \text{ mg/m}^3 \times 3 \text{ h/day} \times 3 \text{ days}$	Sprague-Dawley male rats	SS fumes are cleared from the lung at a slower rate compared to MS fumes
Cohen et al. (1998)	Soluble, insoluble Cr; $360 \mu g/m^3 \times 5 h/day \times 5 days/week \times 2 or 4 weeks alone or in combination with 0.3 ppm O_3$	F344 male rats	Immunomodulatory effects of chromium related to solubility
Erdely et al. (2011a)	GMA-SS; $40 \text{ mg/m}^3 \times 3 \text{ h/day} \times 10 \text{ days}$	C57BL/6J male mice	Pulmonary and systemic induction of type I interferon signaling
Oh et al. (2009)	MMA-SS; 51.4 or 84.6 mg/m ³ × 2 h/day × 30 days plus 30 day recovery	Sprague-Dawley male rats	Transcriptional changes associated with exposure and recovery to MMA-SS
			(Continued

(Continued)



Table 6. (Continuation Author	Welding fume	Animal model	Finding
Oh et al. (2011)	MMA-SS; 51.4 or 84.6 mg/m $^3 \times 2$ h/day \times 30 days then 30 day recovery followed by 2^{nd} exposure of 44.1 or 80.1 mg/m $^3 \times 2$ h/day \times 30 days plus 30 day recovery	Sprague-Dawley male rats	Transcriptional changes with signaling pathways of proposed molecular mechanisms of repeated exposure and recovery to MMA-SS
Rim et al. (2004)	MMA-SS; 107.5 mg/m $^3 \times 2 \text{ h/day} \times 30 \text{ days}$	Sprague-Dawley male rats	Prolonged MMA-SS exposure down-regulated gene expression in peripheral mononuclear cells
Sung et al. (2004)	MMA-SS; 64.8 or 107.8 mg/m 3 × 2 h/day × 60 days	Sprague-Dawley male rats	General pulmonary effects of MMA-SS inhalation
Yang et al. (2009)	MMA-SS; 51.4 or 84.6 mg/m 3 × 2 h/day × 30 days, then 30 day recovery followed by 2^{nd} exposure of 44.1 or 80.1 mg/m 3 × 2 h/day × 30 days plus 30 day recovery	Sprague-Dawley male rats	Recurrent exposures to MMA-SS was more difficult to recover from than a single exposure
Yu et al. (2004)	MMA-SS; 65.6 or 116.8 mg/m 3 × 2 hr/d × 30 d	Sprague-Dawley male rats	MMA-SS induced pulmonary inflammatory and genotoxic responses
Zeidler- Erdely et al. (2011b)	GMA-SS; $40 \text{ mg/m}^3 \times 3 \text{ h/day} \times 6 \text{ or } 10 \text{ days}$	A/J and C57BL/6J male mice	GMA-SS induced a sustained inflammatory response in both strains with no resolution after 1 month
Route of Exposu	re: Intratracheal Instillation		
Antonini and Roberts (2007)	MMA-SS (2 mg/rat), Fe $_2$ O $_3$ (0.82 mg/rat), NiO (0.06 mg/rat), Cr $_2$ Na $_2$ O $_7$ (0.60 mg/rat)	Sprague-Dawley male rats	Cr, not Fe or Ni, in MMA-SS was a primary component for reduced bacterial clearance
Antonini et al. (1996)	SS-Spray, MS-Spray, MS-Pulse; 1 mg/body wt	CD/VAF male rats	SS fumes are cleared from the lung at a slower rate compared to MS fumes
Antonini et al. (2004)	GMA-MS, GMA-SS, MMA-SS; 0.10 or 2.00 mg/rat, inoculation 3 days later with 5×10^3 <i>L. monocytogenes</i>	Sprague-Dawley male rats	MMA-SS, but not GMA-SS or GMA-MS, reduced bacterial clearance after exposure
Taylor et al. (2003)	GMA-MS, GMA-SS, MMA-SS; 2 mg/rat	Sprague-Dawley male rats	Metal composition and soluble and insoluble components affect the toxicity of a given welding fume
Route of Exposu	re: Pharyngeal Aspiration		
Anderson et al. (2007)	MMA-SS; 5–20 mg/kg body wt every 5 days \times 4 doses, immunized 24 h later	B6C3F1 female mice	Reduced pulmonary and systemic humoral immune response despite increased lymphocyte numbers
Erdely et al. (2011b)	GMA-MS, GMA-SS, MMA-SS; 85 μ g × 4 doses	C57BL/6J male mice	Comparison of pulmonary and systemic effects of SS and MS welding fume exposure
Zeidler- Erdely et al. (2008)	GMA-MS, GMA-SS, MMA-SS; 85 μ g × 4 doses	A/J and C57BL/6J male mice	GMA-SS caused prolonged lung inflammation in tumor susceptible A/J mice that may enhance the potential of tumorigenesis
Zeidler- Erdely et al. (2010)	GMA-SS; 85 μg × 4 doses	A/J and C57BL/6J male	GMA-SS caused prolonged lung inflammation in tumor susceptible A/J mice that was associated with a dysregulation of immunomodulatory genes

Abbreviations include gas metal arc (GMA)-stainless steel (SS); GMA-mild steel (MS); manual metal arc (MMA)-SS; soluble (S)-chromium (Cr); nickel (Ni); iron (Fe).

functionality of immune competent cells was associated with altered immune-related cytokine and antibody production as well as impaired resolution of local inflammation. The reduced capacity to eliminate infectious agents results in enhanced pulmonary injury, exacerbating the initial exposure effects and perhaps providing the potential to extend the window of immunosuppression.

Disclaimer

The findings and conclusions of this paper have not been formally disseminated by NIOSH and should not be construed to represent any agency determination or policy.

Declaration of interest

The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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