# US Patent & Trademark Office Patent Public Search | Text View

United States Patent

Kind Code

Date of Patent

Inventor(s)

12385727

August 12, 2025

Yates; Daniel

# Fuzzy interface layer for thermite and primer made from thermite with fuzzy layer

#### Abstract

A layered thermite composite includes alternating layers of metal oxide and reducing metal deposited upon a substrate. A fuzzy interface layer between each metal oxide layer and reducing metal layer includes reducing metal, reducing metal oxide, and metal oxide at least partially intermixed therein. The fuzzy interface layer forms while moving between chambers in a multichamber deposition process which minimizes exposure of the reducing metal to oxygen while moving between chambers. The fuzzy interface layer is not expected to grow in thickness after deposition. The combination of the relative thinness of the fuzzy interface layer as well as the presence of reactants as well as reducing metal oxide maintains the sensitivity of the layered thermite structure to mechanical ignition if the structure is used within a primer.

**Inventors:** Yates; Daniel (Melbourne, FL)

**Applicant: Spectre Primer Technologies, Inc.** (Palm Bay, FL)

Family ID: 1000008751664

Assignee: Spectre Primer Technologies, Inc. (Palm Bay, FL)

Appl. No.: 18/649459

Filed: April 29, 2024

#### **Prior Publication Data**

**Document Identifier**US 20240361113 A1

Publication Date
Oct. 31, 2024

# **Related U.S. Application Data**

us-provisional-application US 63462995 20230429

# **Publication Classification**

Int. Cl.: F42C19/08 (20060101); C06B45/12 (20060101)

U.S. Cl.:

CPC **F42C19/083** (20130101); **C06B45/12** (20130101); **F42C19/0803** (20130101);

F42C19/08 (20130101)

# **Field of Classification Search**

**CPC:** F42C (19/08); F42C (19/083); F42C (19/0803); F42C (19/0819); F42C (19/0823); F42B

(3/10); F42B (5/10); F42B (5/16); C06B (45/12); C06B (45/14); C06B (33/00)

**USPC:** 102/204; 102/470

# **References Cited**

#### **U.S. PATENT DOCUMENTS**

0.5.1711L111 DO	COMENIO			
Patent No.	<b>Issued Date</b>	<b>Patentee Name</b>	U.S. Cl.	CPC
2131352	12/1937	Marsh	N/A	N/A
2239123	12/1940	Stoneking	N/A	N/A
2652775	12/1952	Swanson	N/A	N/A
2995429	12/1960	Williams et al.	N/A	N/A
2995431	12/1960	Bice	N/A	N/A
3122884	12/1963	Grover et al.	N/A	N/A
3155749	12/1963	Rossen et al.	N/A	N/A
3170402	12/1964	Morton et al.	N/A	N/A
3382117	12/1967	Cook	N/A	N/A
3610151	12/1970	Nett	N/A	N/A
3668872	12/1971	Camp et al.	N/A	N/A
3711344	12/1972	Pierce	N/A	N/A
3715248	12/1972	Swotinsky et al.	N/A	N/A
3725516	12/1972	Kaufman	N/A	N/A
3808061	12/1973	Pierce	N/A	N/A
3896731	12/1974	Comfort et al.	N/A	N/A
3905846	12/1974	Berta	N/A	N/A
3938440	12/1975	Dooley et al.	N/A	N/A
3956890	12/1975	Davis	N/A	N/A
3961576	12/1975	Montgomery, Jr.	N/A	N/A
3962865	12/1975	McCone, Jr.	N/A	N/A
3995559	12/1975	Bice	N/A	N/A
4013743	12/1976	Blasche, Jr	N/A	N/A
4115999	12/1977	Diebold	N/A	N/A
4475461	12/1983	Durrell	N/A	N/A
4615270	12/1985	Bell	N/A	N/A
4651254	12/1986	Brede et al.	N/A	N/A
4756251	12/1987	Hightower, Jr. et al.	N/A	N/A

4823699	12/1988	Farinacci	N/A	N/A
4823701	12/1988	Wilhelm	N/A	N/A
4875948	12/1988	Verneker	N/A	N/A
4963203	12/1989	Halcomb et al.	N/A	N/A
4996922	12/1990	Halcomb	N/A	N/A
5030301	12/1990	Stout et al.	N/A	N/A
5076868	12/1990	Doll et al.	N/A	N/A
5080017	12/1991	Asikainen	N/A	N/A
5237927	12/1992	Gonzalez	N/A	N/A
5266132	12/1992	Danen et al.	N/A	N/A
5320043	12/1993	Andre et al.	N/A	N/A
5322018	12/1993	Hadden	N/A	N/A
5363768	12/1993	Solberg	N/A	N/A
5378499	12/1994	Martin et al.	N/A	N/A
5589661	12/1995	Menke et al.	N/A	N/A
5721392	12/1997	Chan	N/A	N/A
5773748	12/1997	Makowiecki	N/A	N/A
5801325	12/1997	Willer et al.	N/A	N/A
5817970	12/1997	Feierlein	N/A	N/A
5854439	12/1997	Almstrom et al.	N/A	N/A
5879519	12/1998	Seeser et al.	N/A	N/A
6090756	12/1999	Brown	N/A	N/A
6158348	12/1999	Campoli	N/A	N/A
6176950	12/2000	Wood et al.	N/A	N/A
6183569	12/2000	Mohler	N/A	N/A
6328856	12/2000	Seeser et al.	N/A	N/A
6334394	12/2001	Zimmerman et al.	N/A	N/A
6363853	12/2001	Rohr	N/A	N/A
6364975	12/2001	Fleming et al.	N/A	N/A
6599379	12/2002	Hiskey et al.	N/A	N/A
6627013	12/2002	Carter, Jr. et al.	N/A	N/A
6679960	12/2003	Jones	N/A	N/A
6692655	12/2003	Martins et al.	N/A	N/A
6712917	12/2003	Gash et al.	N/A	N/A
6740180	12/2003	Cesaroni	N/A	N/A
6805832	12/2003	Mohler et al.	N/A	N/A
6843868	12/2004	Fawls et al.	N/A	N/A
6962112	12/2004	Kern	N/A	N/A
7770380	12/2009	Dulligan et al.	N/A	N/A
7886668	12/2010	Hugus et al. Mohler	N/A	N/A
7896988 7918163	12/2010 12/2010		N/A N/A	N/A N/A
7916163 7955451	12/2010	Dahlberg Hugus et al.	N/A N/A	N/A N/A
7958823	12/2010	Sawka	N/A N/A	N/A N/A
7998290	12/2010	Sheridan et al.	N/A N/A	N/A N/A
8202377	12/2010	Erickson et al.	N/A N/A	N/A N/A
8298358	12/2011	Coffey et al.	N/A N/A	N/A N/A
8454769	12/2011	Erickson et al.	N/A N/A	N/A N/A
8465608	12/2012	Coffey et al.	N/A	N/A
8505427	12/2012	Wilson et al.	N/A	N/A
0000-T4/	1 <i>4</i> /4014	Willoui Ct ui.	1 1/ 1 1	1 1/ / 1

8524018	12/2012	Busky et al.	N/A	N/A
8544387	12/2012	Dahlberg	N/A	N/A
8591676	12/2012	Coffey et al.	N/A	N/A
8641842	12/2013	Hafner et al.	N/A	N/A
8758580	12/2013	DeVito	N/A	N/A
9255775	12/2015	Rubin	N/A	N/A
9464874	12/2015	Mohler et al.	N/A	N/A
9709366	12/2016	Mohler et al.	N/A	N/A
9816792	12/2016	Mohler et al.	N/A	N/A
10254090	12/2018	Mohler et al.	N/A	N/A
10415938	12/2018	Mohler et al.	N/A	N/A
10882799	12/2020	Coffey et al.	N/A	N/A
10989510	12/2020	Mohler et al.	N/A	N/A
11112222	12/2020	Coffey et al.	N/A	N/A
11650037	12/2022	Yates	102/430	C06B
				33/00
2002/0092438	12/2001	Makowiecki	N/A	N/A
2006/0011276	12/2005	Grix et al.	N/A	N/A
2006/0278119	12/2005	Shilliday et al.	N/A	N/A
2007/0099335	12/2006	Gangopadhyay et	N/A	N/A
		al.		
2007/0169862	12/2006	Hugus et al.	N/A	N/A
2007/0272112	12/2006	Nielson et al.	N/A	N/A
2008/0028922	12/2007	Wilson	N/A	N/A
2008/0047453 2008/0134924	12/2007 12/2007	Dahlberg Sawka	N/A N/A	N/A N/A
2006/0134924	12/2007	Mohler	N/A N/A	N/A N/A
2009/0178741	12/2008	Xun et al.	N/A N/A	N/A N/A
2010/0193093	12/2009	Coffey et al.	N/A	N/A N/A
2010/0153053	12/2009	Hugus et al.	N/A	N/A
2010/0282115	12/2009	Sheridan et al.	N/A	N/A
2011/0067789	12/2010	Grix et al.	N/A	N/A
2011/0007703	12/2010	Sawka et al.	N/A	N/A
2011/0308416	12/2010	Bar et al.	N/A	N/A
2012/0103479	12/2011	Katzakian et al.	N/A	N/A
2012/0132096	12/2011	Chin et al.	N/A	N/A
2013/0305950	12/2012	Coffman, II	N/A	N/A
2014/0237879	12/2013	Mirabile	N/A	N/A
2016/0102030	12/2015	Coffey et al.	N/A	N/A
2016/0216095	12/2015	Rami	N/A	N/A
2019/0128656	12/2018	Mohler et al.	N/A	N/A
2020/0232772	12/2019	Coffey et al.	N/A	N/A
2020/0400415	12/2019	Mohler et al.	N/A	N/A
2022/0260353	12/2021	Yates	N/A	N/A
2024/0238956	12/2023	Mohler	N/A	B25C
202 <del>1</del> /0230330	14/4040	IVIOIIICI	1 1/ 1 1	1/143
FOREIGN PATENT DOCUMENTS				

Application Date Patent No. Country CPC

0122012	12/1983	EP	N/A
328257	12/1988	EP	N/A
885409	12/1960	GB	N/A
944442	12/1962	GB	N/A
987332	12/1964	GB	N/A
994184	12/1964	GB	N/A
2005121055	12/2004	WO	N/A
WO-2025085106	12/2024	WO	F42C 19/0803

#### OTHER PUBLICATIONS

International Search Report for PCT/US2024/026858 dated Apr. 30, 2025. cited by applicant International Written Opinion for PCT/US2024/026858 dated Apr. 30, 2025. cited by applicant Zirconium, Wikipedia, https://en.wikipedia.org/wiki/Zirconium, dated at least as early as Jan. 1, 2021. cited by applicant

Zirconium Carbide, Wikipedia, https://en.wikipedia.org/wiki/Zirconium\_carbide, dated at least as early as Jan. 1, 2021. cited by applicant

K. Woll et al., The utilization of metal/metal oxide core shell powders to enhance the reactivity of diluted thermite mixtures, Combustion and Flame 167 (2016) 259-267. cited by applicant A. H. Kinsey et al., Gas Suppression via Copper Interlayers in Magnetron Sputtered Al—Cu2O Multilayers, ACS Applied Materials & Interfaces, 2017, 22026-22036. cited by applicant Aluminum Carbide, Wikipedia, https://en.wikipedia.org/wiki/Aluminium\_carbide, dated at least as early as Jan. 1, 2021. cited by applicant

K. J. Blobaum et al., Deposition and characterization of a self-propagating CuOx/Al thermite reaction in a multilayer foil geometry, J. Appl. Phys., vol. 94, No. 5, 2003. cited by applicant Carbide, Wikipedia, https://en.wikipedia.org/wiki/Carbide, dated at least as early as Jan. 1, 2021. cited by applicant

Ceramic, Wikipedia, https://en.wikipedia.org/wiki/Ceramic, dated at least as early as Jan. 1, 2021. cited by applicant

Detonator, Wikipedia, https://en.wikipedia.org/wiki/Detonator, dated at least as early as Jan. 1, 2021. cited by applicant

High Power Impulse Magnetron Sputtering, Wikipedia, https://en.wikipedia.org/wiki/High-power\_impulse\_magnetron\_sputtering, dated at least as early as Jan. 1, 2021. cited by applicant J. L. Gottfried, Improving the Explosive Performance of Aluminum Nanoparticles with aluminum lodate Hexahydrate (AIH), 2018, https://www.nature.com/articles/s41598-018-26390-9. cited by applicant

G. C. Egan et al., Probing the Reaction Dynamics of Thermite Nanolaminates, J. Phys. Chem. 119, 2015, 20401-20408. cited by applicant

Thermite, Wikipedia, https://en.wikipedia.org/wiki/Thermite, dated at least as early as Jan. 1, 2021. cited by applicant

Titanium Carbide, Wikipedia, https://en.wikipedia.org/wiki/Titanium\_carbide, dated at least as early as Jan. 1, 2021. cited by applicant

Versatile Reactive Sputtering Batch Drum Coater With Auxiliary Plasma, Deposition Sciences, Inc., 2004. cited by applicant

Primary Examiner: Bergin; James S

Attorney, Agent or Firm: Lang Patent Law LLC

# **Background/Summary**

CROSS REFERENCE TO RELATED APPLICATION (1) This application claims the benefit of U.S. provisional patent application Ser. No. 63/462,995, which was filed on Apr. 29, 2023, and entitled "Fuzzy Interface Layer For Thermite And Primer Made From Thermite With Fuzzy Layer."

#### **TECHNICAL FIELD**

- (1) The present invention relates to primers for firearms and other munitions. More specifically, a primer made from layered metal oxide and reducing metal with a fuzzy interface layer between the reducing metal and metal oxide is provided BACKGROUND INFORMATION
- (2) Cartridges for firearms, as well as other munitions such as larger projectile cartridges and explosives are often ignited by a primer. Presently available primers and detonators are made from a copper or brass alloy cup with a brass anvil and containing lead azide or lead styphnate. When the base of the cup is struck by a firing pin, the priming compound is crushed between the cup's base and the anvil, igniting the primer charge. The burning primer then ignites another flammable substance such as smokeless powder, explosive substances, etc. Lead azide and lead styphnate are hazardous due to their toxicity as well as their highly explosive nature. Additionally, present manufacturing methods are very labor-intensive, with the necessary manual processes raising costs, causing greater difficulty in maintaining quality control.
- (3) Energetic materials such as thermite are presently used when highly exothermic reactions are needed. Uses include cutting, welding, purification of metal ores, and enhancing the effects of high explosives. A thermite reaction occurs between a metal oxide and a reducing metal. Examples of metal oxides include La.sub.2O.sub.3, AgO, ThO.sub.2, SrO, ZrO.sub.2, UO.sub.2, BaO, CeO.sub.2, B.sub.2O.sub.3, SiO.sub.2, V.sub.2O.sub.5, Ta.sub.2O.sub.5, NiO, Ni.sub.2O.sub.3, Cr.sub.2O.sub.3, MoO.sub.3, P.sub.2O.sub.5, SnO.sub.2, WO.sub.2, WO.sub.3, Fe.sub.3O.sub.4, CoO, Co.sub.3O.sub.4, Sb.sub.2O.sub.3, PbO, Fe.sub.2O.sub.3, Bi.sub.2O.sub.3, MnO.sub.2, Cu.sub.2O, and CuO. Example reducing metals include Al, Zr, Th, Ca, Mg, U, B, Ce, Be, Ti, Ta, Hf, and La. The reducing metal may also be in the form of an alloy or intermetallic compound of the above-listed metals.
- (4) The reducing metals used within energetic materials will oxidize if exposed to oxygen or to water vapor. Prior art thermite structures attempted to balance the increased ignition rate of having the reducing metal and metal oxide more finely divided with the reduced energy density which results from a greater portion of the reducing metal oxidizing prior to ignition. For example, in a layered thermite composite, thinner layers within a composite having the same overall thickness will typically have a faster ignition rate but a lower energy density than a composite having thicker layers of reactants. One prior art reference, US 2002/0092438, attempted to characterize the gradual increase of the size of the interface layer over time as a positive feature, claiming it could be used to limit the lifespan of a primer for ammunition. Minimizing the thickness of the reducing metal oxide interface layer, and/or changing the composition of the interface layer can permit increased ignition rates without compromising energy density. Particularly when such a thermite composite is deposited on a malleable substrate, a thin interface layer and/or an interface layer which includes reactant atoms as well as reducing metal oxide atoms at least partially intermixed therein could aid in susceptibility to mechanical ignition.
- (5) Accordingly, there is a need for a primer made from materials that do not share the toxicity of lead. There is an additional need for a layered thermite composite having interface layers between reactants which do not significantly impact energy density or susceptibility to mechanical ignition. There is a further need for a primer made from materials that lend themselves to automated processes, as well as processes which protect the reducing metal from oxidizing during and after deposition. Another need exists for a primer made from energetic materials that lends itself to

ignition through a strike by a firing pin, but which otherwise benefits from the stability of thermite. SUMMARY

- (6) The above needs are met by a layered thermite composite. The thermite composite comprises a substrate having a deposition surface and a rear surface. Alternating layers of metal oxide and reducing metal are deposited upon the substrate. The alternating layers of metal oxide and reducing metal are structured to react with each other in response to an impact applied to the rear surface of the substrate. The thermite composite further comprises a fuzzy interface layer disposed between each metal oxide layer and reducing metal layer. The fuzzy interface layer contains reducing metal, reducing metal oxide, and metal oxide which are at least partially mixed together.
- (7) The above needs are further met by a primer. The primer comprises a substrate having a deposition surface and a rear surface. The primer further has alternating layers of metal oxide and reducing metal deposited upon the substrate. The alternating layers of metal oxide and reducing metal are structured to react with each other in response to an impact applied to the rear surface of the substrate. The primer also comprises a fuzzy interface layer disposed between each metal oxide layer and reducing metal layer. The fuzzy interface layer contains reducing metal, reducing metal oxide, and metal oxide which are at least partially mixed together.
- (8) The above needs are additionally met by a cartridge for a firearm. The cartridge comprises a casing having a front end, a back end, and a hollow interior. A bullet is secured within the front end of the casing. Aa propellant is disposed within the hollow interior. A primer is secured within the back end of the casing. The primer is in communication with the propellant. The primer comprises a substrate having a deposition surface and a rear surface. The primer further has alternating layers of metal oxide and reducing metal deposited upon the substrate. The alternating layers of metal oxide and reducing metal are structured to react with each other in response to an impact applied to the rear surface of the substrate. The primer also has a fuzzy interface layer disposed between each metal oxide layer and reducing metal layer. The fuzzy interface layer contains reducing metal, reducing metal oxide, and metal oxide which are at least partially mixed together.
- (9) These and other aspects of the invention will become more apparent through the following description and drawings.

# Description

#### BRIEF DESCRIPTION OF THE DRAWINGS

- (1) FIG.  ${\bf 1}$  a sectional, side elevational view of an example of a layered thermite composite according to the present invention.
- (2) FIG. **2** is a sectional, side elevational view of a layered thermite composite, a carbide-containing ceramic layer, and passivation coating of a primer.
- (3) FIG. **3** is a sectional, side elevational view of an alternative layered thermite composite, a pair of carbide-containing ceramic layers, and passivation coating of a primer.
- (4) FIG. **4** is a sectional, side elevational view of another alternative layered thermite composite, a carbide-containing ceramic layer, and passivation coating of a primer.
- (5) FIG. **5** is a sectional, side elevational view of a fuzzy interface layer between a metal oxide layer and a reducing metal layer of the thermite structure of FIG. **1**.
- (6) FIG. **6**A is a sectional, side elevational view of the box A in FIG. **5**.
- (7) FIG. **6**B is a sectional, side elevational view of the box A in FIG. **5**, showing the oxygen content within the box A of FIG. **5**.
- (8) FIG. **6**C is a sectional, side elevational view of the box A in FIG. **5**, showing the aluminum content within the box A of FIG. **5**.
- (9) FIG. **6**D is a sectional, side elevational view of the box A in FIG. **5**, showing the copper content within the box A of FIG. **5**.

- (10) FIG. **7** is a graph showing the atomic percent of aluminum, oxygen, and copper with respect to position within the fuzzy interface of FIGS. **5-6**D.
- (11) FIG. **8** is a side elevational, cross sectional view of a cup for use with a primer material of FIGS. **1-3**.
- (12) FIG. **9** is a side elevational, cross sectional view of a cartridge using a primer cup of FIG. **4**.
- (13) Like reference characters denote like elements throughout the drawings.

#### **DETAILED DESCRIPTION**

- (14) Referring to FIGS. **1-4**, a layered thermite composite **10** is shown. The layered thermite composite is particularly useful as a portion of a primer **14**, as well as for other uses. The layered thermite composite **10** is deposited upon a substrate **12**. The layered thermite composite **10** may include one or more carbide-containing ceramic layer(s) **16** within the layered thermite coating **14**, and includes a passivation coating **18**.
- (15) If the thermite composite **10** is within a primer **14**, then the substrate **12** in the illustrated example is a malleable disk, made from a material such as brass, copper, soft steel, and/or stainless steel, having a deposition surface **19** upon which the layered thermite coating **10** is deposited, and a rear surface **21** (FIG. **4**). The substrate **12** is a sufficiently thin and malleable so that a firing pin strike to the rear surface **21** will ignite the layered thermite coating **10** and carbide-containing ceramic layer(s) **16** as described below, but is sufficiently thick for ease of manufacturing the primer **14** as well as securing a primer **14** within a cartridge case, munition, modified primer cup, or other location as described below. A preferred substrate thickness is about 0.005 inch to about 0.1 inch, and is more preferably about 0.01 to about 0.025 inch.
- (16) The layered thermite coating **14** includes alternating layers of metal oxide and reducing metal (with only a small number of layers illustrated for clarity). Examples of metal oxides include La.sub.2O.sub.3, AgO, ThO.sub.2, SrO, ZrO.sub.2, UO.sub.2, BaO, CeO.sub.2, B.sub.2O.sub.3, SiO.sub.2, V.sub.2O.sub.5, Ta.sub.2O.sub.5, NiO, Ni.sub.2O.sub.3, Cr.sub.2O.sub.3, MoO.sub.3, P.sub.2O.sub.5, SnO.sub.2, WO.sub.2, WO.sub.3, Fe.sub.3O.sub.4, CoO, Co.sub.3O.sub.4, Sb.sub.2O.sub.3, PbO, Fe.sub.2O.sub.3, Bi.sub.2O.sub.3, MnO.sub.2, Cu.sub.2O, and CuO. Example reducing metals include Al, Zr, Th, Ca, Mg, U, B, Ce, Be, Ti, Ta, Hf, and La. The metal oxide and reducing metal are preferably selected to resist abrasion or other damage to a barrel of a firearm with which a cartridge containing the primer is used by avoiding reaction products which could potentially cause such damage. One example of such a combination of metal oxide and reducing metal is cupric oxide and magnesium.
- (17) The thickness of each metal oxide layer and reducing metal layer are determined to ensure that the proportions of metal oxide and reducing metal are such so that both will be substantially consumed by the exothermic reaction. As one example, in the case of a metal oxide layer **20** made from CuO and reducing metal layer **22** made from Al (FIG. **1**), the chemical reaction is 3CuO+2Al.fwdarw.3Cu+Al.sub.2O.sub.3+heat. The reaction therefore requires 3 moles of CuO, weighing 79.5454 grams/mole, for every 2 moles of Al, weighing 26.98154 grams/mole. CuO has a density of 6.315 g/cm.sup.3, and aluminum has a density of 2.70 g/cm.sup.3. Therefore, the volume of CuO required for every 3 moles is 37.788 cm.sup.3. Similarly, the volume of Al required for every 2 moles is 19.986 cm.sup.3. Therefore, within the illustrated example of a composite layer **16**, the metal oxide **12** is about twice as thick as the reducing metal **14**.
- (18) As another example, in the case of a metal oxide layer **20** made from CuO and reducing metal layer **22** made from Mg, the chemical reaction is CuO+Mg.fwdarw.Cu+MgO+heat. The reaction therefore requires one mole of CuO, weighing 79.5454 grams/mole, for every one mole of Mg, weighing 24.305 grams/mole. CuO has a density of 6.315 g/cm.sup.3, and magnesium has a density of 1.74 g/cm.sup.3. Therefore, the volume of CuO required for every mole is 12.596 cm.sup.3. Similarly, the volume of Mg required for every mole is 13.968 cm.sup.3. Therefore, within the illustrated example, each layer of metal oxide is about the same thickness or slightly thinner than the corresponding layer of reducing metal. If other metal oxides and reducing metals

are selected, then the relative thickness of the metal oxide and reducing metal can be similarly determined.

(19) Referring to FIGS. **5-6**D, the illustrated example of the thermite structure **10** also includes a fuzzy interface layer **24** between each reducing metal layer **16** and adjacent metal oxide layer **14**. As used herein, a fuzzy interface layer **24** is an interface between a reducing metal layer **16** and a metal oxide layer 14, with the interface layer 24 containing metal oxide 20, reducing metal 22, and reducing metal oxide, all of which are at least partially intermixed to form a gradient structure within the interface layer 24. (Due to the intermixing of materials at the interface layer 24, reference characters **14** and **16** refer to metal oxide layers and reducing metal layers, respectively, while reference characters **20** and **22** refer to metal oxide and reducing metal, respectively, regardless of whether those materials are within a layer **14** or **16**.) Although the approximate thickness of the fuzzy interface layer **24** is about 2 nm to about 5 nm, the fuzzy interface layer **24** does not have precise boundaries. Instead, the amount of each material present in and around the fuzzy interface layer **24** will be a gradient with respect to proximity to either the reducing metal layer **16** or metal oxide layer **14**. As shown in the example of FIG. **5**, moving from the bottom of the image to the top, the material transitions from aluminum 22 within the reducing metal layer 16, to a combination of aluminum 22, aluminum oxide, and cupric oxide 20 as the interface layer 24 within the approximate center of the image is reached. Continuing upward in the image, the material again transitions from the mixture of aluminum 22, aluminum oxide, and cupric oxide 20 in the interface layer **24** to simply cupric oxide in the metal oxide layer **14**. (20) A similar gradient pattern is shown in the examples of FIGS. **6**A-**6**D, with FIG. **6**A simply showing the detail of box A in FIG. 5. FIG. 6B focuses on the oxygen 26 present within Box A. In FIG. **6**B, the aluminum layer **16** contains no oxygen **26**, but oxygen in relatively high concentration (appearing as a lighter shade) is present above the aluminum **28** within the interface **24**. Cupric oxide **20** is present above the oxygen **26** in relatively high concentration within the layer **24** as well as the layer **16**. FIG. **6**C focuses on aluminum **28** (shown in a lighter shade), showing the transition between pure aluminum within the layer 16, to a mixture of aluminum 28 and other elements in the interface **24**, and no aluminum **28** in the layer **14**. Similarly, FIG. **6**D focuses on the copper **30** (shown in a lighter shade), showing no copper **30** in the layer **16**, some copper **30** in the interface **24**, and a large percentage of copper **30** in the layer **14**. The atomic percent of each element within box A of FIG. 5 is also graphically illustrated in FIG. 7, with the left side 32 of FIG. 7 corresponding to the bottom of the images of FIGS. 5-6D, and the right side 34 of FIG. 7 corresponding to the top of the images **5-6**D. Line **36** shows aluminum, line **38** shows oxygen, and line **40** shows copper. As shown in FIG. **7** the atomic percent aluminum is maximized on the left side **32**, decreasing towards zero at some position within the layer **24**. The atomic percentage of oxygen (within aluminum oxide and cupric oxide) is very small in the layer **16**, but becomes high in the layer **24**, and decreases to a stable percentage within the layer **14**. Copper (in the form of cupric oxide) is absent from the layer **16**, but increases as the layer **24** is entered, increasing throughout the layer **24** until stabilizing in the layer **14**. Although other examples of fuzzy interface layers will follow similar patterns, variation will occur in the location of the transitions, atomic percent of each element at various locations, and the overall thickness of the fuzzy interface layer. (21) The interface layer **24** forms between completion of depositing one layer of reducing metal **16** or metal oxide **14** and the beginning of deposition of the next layer of reducing metal **16** or metal oxide **14**. Prior art interface layers would form as the surface of the reducing metal oxidized from exposure to atmospheric oxygen or water vapor, and were thus composed of reducing metal oxide. The fuzzy interface layer described herein is formed by a process (described in greater detail below) which permits rapid transitions from depositing one type of layer to depositing the other type of layer, permitting only a limited amount of reducing metal oxide to form during the transition between depositing reducing metal and depositing metal oxide. The resulting interface layer is therefore a gradient structure of metal oxide, reducing metal, and reducing metal oxide

rather than pure reducing metal oxide.

- (22) A layered thermite composite **10** can be made using a deposition system on which the substrate is secured to a substrate support which is movable between a plurality of deposition chambers. As one example, the substrate support may use a substrate support in the form of a rotating drum having a surface on which the substrates are secured, and includes a plurality of deposition chambers positioned around the drum. Such systems are described in the following patents or published applications, the entire disclosure of all of which are expressly incorporated herein by reference: U.S. Pat. No. 8,758,580, which was issued to R. DeVito on Jun. 24, 2014; U.S. Pat. No. 5,879,519, which was issued to J. W. Seeser et al. on Mar. 9, 1999; EP 0,328,257, which was invented by M. A. Scobey et al. and published on Aug. 16, 1989, and U.S. Pat. No. 6,328,856, which was issued to J. W. Seeser et al. on Dec. 11, 2001. The use of a rotating drum system permits the substrates to be rapidly transferred between different chambers for deposition of different layers made from different materials. In one example, some chamber(s) will be used to deposit the reducing metal, other chamber(s) will be used to deposit the metal oxide, and still other chamber(s) may be used to deposit the carbide-containing ceramic (if a primer is the intended result). In a four chamber system, other chambers may be used to deposit the adhesion layers above and below the carbide-containing ceramic. One example may utilize between two and four chambers, with two targets per chamber. The atmospheric conditions within each chamber are maintained and isolated from other portions of the system by baffles which extend close to the drum while maintaining separation from the substrates. Substrates may thereby be moved between chambers by rotating the drum upon which the substrates are located while maintaining the correct pressure and atmospheric conditions of each chamber throughout the process of depositing multiple layers. During the process of depositing metal oxide and reducing metal, the individual chambers will run continuously, and the drum or other substrate carrier will rotate or otherwise move continuously to move the substrates through the appropriate chambers at the appropriate rate. Additionally, the pressure of an inert gas, for example, argon in the chamber utilized to deposit reducing metal may be greater than the pressure in the chamber utilized to deposit metal oxide, thus resisting the entry of oxygen into the reducing metal chamber. The need to pump down each chamber between layers of different material is thus avoided, speeding and simplifying the deposition process. (23) Prior art manufacturing methods typically required several minutes of deposition time for each of the reducing metal or metal oxide layers, with multiple minutes of additional time required to switch from depositing one material to depositing the other material. The above-described process permits each layer to be deposited in a time of, for example, about 15 seconds. Transitioning from one chamber to the next chamber can be accomplished in a time of, for example, about 2 seconds. The manufacturing process is thus significantly faster, as well as providing very little time for interface layers having undesirable characteristics to form. Without being bound by any particular theory, it is believed that the oxygen which reacts with the reducing metal during transitions between chambers is atmospheric oxygen and/or oxygen from the deposition of the metal oxide rather than oxygen from water vapor. Again without being bound by any particular theory, it is believed that interface layers formed by reactions with water vapor are more likely to grow over time through additional reaction with and oxidation of the reducing metal. Interfaces formed by reactions with atmospheric oxygen and/or oxygen from the deposition of metal oxide will resist additional metal oxide formation once the interface is covered by the next layer of reactant. Because the fuzzy interface layer **24** will not grow over time, and because the fuzzy interface region includes not only reducing metal oxide but also metal oxide and reducing metal, the metal oxide and reducing metal remain in sufficiently close proximity to each other so that they can be ignited electrically or mechanically when desired.
- (24) If the thermite structure is intended for use as a primer **14**, then some examples of the primer **14** may include elements which will either facilitate ignition and/or facilitate carrying the ignition to a propellant within a cartridge casing or to another ignitable material, for example, a fuse which

is intended for ignition by the primer.

- (25) The illustrated example in FIGS. 2 and 3 of a layered thermite coating 14 is divided into an initial ignition portion **42** that is deposited directly onto the substrate **12**, and a secondary ignition portion **44** that is deposited onto the initial ignition portion **42**. The illustrated example of the initial ignition portion **42** includes layers of metal oxide **46** and reducing metal **48** that are thinner than the layers of metal oxide **50** and reducing metal **52** within the secondary ignition portion **44**. In the illustrated example, each metal oxide **46** and reducing metal **48** pair of layers are preferably between about 20 nm and about 100 nm thick, with the illustrated example having pairs of layers that are about 84 nm thick. In the illustrated example, each pair of metal oxide 50 and reducing metal **52** layers are thicker than about 100 nm thick. Thinner layers result in more rapid burning and easier ignition, while thicker layers provide a slower burn rate. The thinner layers **46**, **48** within the initial ignition portion **42** are more sensitive to physical impacts, thereby facilitating ignition in response to a firing pin strike to the rear surface **21** of the substrate **12**, and ignite the secondary ignition portion **44**. The thicker layers **50**, **52** within the secondary ignition portion **44** burn more slowly, enhancing the reliability of the ignition of the smokeless powder, explosive, or other desired ignitable substance. The total thickness of the illustrated examples of the layered thermite coating **10** is between about 25 µm and about 1,000 µm.
- (26) The illustrated example of the thermite coating **10** in FIGS. **2** and **3** shows a generally uniform thickness for all layers **46**, **48** within the initial ignition portion **42**. Similarly, a generally uniform thickness is shown within the layers **50**, **52** within the secondary ignition portion **44**. Other examples may include metal oxide and reducing metal layers having differing thicknesses. For example, FIG. **4** shows a primer composition **10** having thermite layers that increase generally proportionally with the distance of the layer from the substrate 12 (with only a small number of layers shown for clarity). Layers 54 and 56, which are close to the substrate 12, have a smaller thickness, for example, between about 20 nm and about 100 nm thick. Layers 58 and 60 have increased thickness. Layers **62** and **64**, farther still from the substrate **12**, have greater thickness than layers **58** and **60**. Layers **66** and **68**, adjacent to the passivation coating **18** and farthest from the substrate 12, are the thickest layers, and are thicker than about 100 nm thick. As before, the total thickness of the illustrated examples of the layered thermite coating **10** is between about 25 μm and about 1,000 μm. Such a thermite coating **10** would provide essentially the same advantage of rapid ignition close to the substrate 12, and relatively slower burning farther from the substrate **12** and closer to the smokeless powder, explosive, or other ignitable substance. With such gradually increasing thickness, a clear boundary between an initial ignition portion and secondary ignition portion may not exist, and a definite boundary is not essential to the functioning of the invention. (27) As another example, all layers of metal oxide and reducing metal may be less than about 100 nm thick, and the time required to consume all layers of metal oxide and reducing metal may be increased sufficiently to ignite conventional propellants and explosives by simply increasing the number of layers of metal oxide and reducing metal.
- (28) Other examples of the layered thermite coating **14** may include layers **46**, **48**, **50**, **52**, or layers **54**, **56**, **58**, **60**, **62**, **64**, **66**, **68**, that are deposited under different temperatures, so that each layer is deposited under a temperature which is either sufficiently higher or sufficiently lower than the adjacent layers to induce thermal expansion and contraction stresses within the layered thermite coating **10** once temperature is equalized within the layered thermite coating. Such expansion and contraction stresses are anticipated to result in increased sensitivity to ignition through a physical impact.
- (29) A passivation layer **18** covers the layered thermite coating **14**, protecting the metal oxide and reducing metal within the layered thermite coating **14**. One example of a passivation layer **18** is silicon nitride. Alternative passivation layers **18** can be made from reactive metals that self-passivate, for example, aluminum or chromium. When oxide forms on the surface of such metals, the oxide is self-sealing, so that oxide formation stops once the exposed surface of the metal is

completely covered with oxide.

- (30) If the layered thermite composite is used for a primer for firearms or other munitions, then the layered structure may include one or more carbide-containing ceramic layer(s). The carbide-containing ceramic layer(s) **16** are disposed within the thermite layers **10**. In the illustrated examples, one carbide-containing ceramic layers **16** is disposed about ⅓ of the distance to the top of the thermite coating **10**. In other examples, a carbide-containing ceramic layer **16** may be located elsewhere in the thermite coating **10**, such as a lower portion, a central portion, the top, the bottom, or elsewhere in the upper portion of the thermite coating **10**. Some examples may include a plurality of layers carbide-containing ceramic layers **16** which are located in different positions throughout the thermite coating **10**. Although one or two layers are illustrated, three or more layers may be utilized. The thickness of the carbide-containing ceramic layer(s) **16** is thicker than the metal oxide or reducing metal layers, and in the illustrated example is between about 100 nm and about 2 μm thick. Other examples of the carbide-containing ceramic layer(s) **16** may be between about 500 nm and about 1 μm thick.
- (31) Carbide-containing ceramics are selected for their propensity, when ignited by ignition of the adjacent reducing metal and metal oxide, to project relatively large (as compared to the thermite reaction products) particles into the propellant of a firearm cartridge or other ignitable or detonatable material. Examples include ceramics such as zirconium carbide, titanium carbide, or silicon carbide, as well as aluminum carbide (which is a metal-ceramic composite but will be considered to be a carbide-containing ceramic herein), and combinations thereof. If more than one carbide-containing ceramic layer is present, then the different carbide-containing ceramic layers may be composed of the same carbide-containing ceramic, or different carbide-containing ceramics. Ignition of these carbides (or other suitable carbides) will result in the formation of carbon dioxide through the reaction with oxygen from the cupric oxide. This gas production will aid in propelling the reaction products of the thermite as well as the reaction products of the carbide-containing ceramic into the propellant or other ignitable or detonatable material. The large, hot particles resulting from the reaction of the carbide-containing ceramic with oxygen will burn for a sufficient period of time to ensure reliable ignition of the propellant or other ignitable or detonatable material.
- (32) Some examples of the layered thermite composite **10** may include an adhesion layer **17** above and below each carbide-containing ceramic layer **16**. In the illustrated example, the adhesion layers 17 are made from titanium or chromium. Nickel may also be used as an adhesion layer in some examples. The illustrated examples of the adhesion layers **17** are about 5 nm to about 10 nm thick. (33) FIG. **8** illustrates an example of a primer **70** utilizing the layered thermite composite **10**. The illustrated example of the substrate 12 is a disk having an upper surface 72 defining a recess 74 in which the deposition surface **19** is located. The edge of the disk **12** includes a larger diameter portion **76** and a smaller diameter portion **78**, forming a ledge **80** therebetween. The primer composite **10** is deposited on the surface **19** within the recess **74** as described above. The disk (substrate) **12** is then placed within a cup **82** to form a complete primer. The cup **82** includes a sidewall **84** having an upper end **86** and a lower end **88**. The lower end **88** includes an inward projection **90** that is dimensioned and configured to abut the ledge **80** and a smaller diameter **78** of the disc **12**. When the disc **12** is inserted into the cup **82** through the upper end **86**, and then placed in position against the lower end **88**, passage of the disc **12** out of the bottom end **88** of the cup **82** is thus resisted. The disc **12** may then be retained in the cup **82** by the inward projections **92** which engage the top surface 72 of the disc. The inward projections 92 may be formed by punching inward against the outer portion of the wall **84** to form depressions **94**, thus creating a projection **92**. Some examples may also, or alternatively retain the disc **12** within the cup **82** utilizing an adhesive.
- (34) Referring to FIG. **9**, the primer **70** may then be placed within a conventional firearm cartridge **96**. The cartridges **96** includes a casing **98** having a standard configuration. The casing **98** includes

a front end **100** that is structured to retain a bullet **102** therein. The casing **98** also includes a back end **104** having a groove **106** and rim **108** to assist with extraction of the cartridge **96**. A propellant **110** within the hollow central portion **112** of the casing **98**. The back end **104** of the casing **98** defines a primer pocket **114** and a flash hole **116** extending between the primer pocket **114** and hollow central portion **112**. Striking the surface **21** with a firing pin ignites the priming compound **10**, driving reaction products through the flash hole **116** and into the propellant **110** discharge the bullet **102**.

- (35) As another example, the layered thermite composite **10** can be used as the deposited ignitable material within the primer disclosed within US 2020/0400415, which was invented by Timothy Mohler and Daniel Yates and published on Dec. 24, 2020, the entire disclosure of which is expressly incorporated herein by reference.
- (36) Although the illustrated examples are for a firearm cartridge, a primer made with the layered thermite composite **10** can be used for a larger projectile cartridge such as those for artillery, or for other munitions such as hand grenades and other explosives that utilize a primer as part of their detonation mechanism.
- (37) The present invention therefore provides a primer made from materials that do not have the toxicity or other safety issues of conventional primers. The primers are easily and inexpensively manufactured by methods that lend themselves to automation. The multi-chamber deposition process eliminate the need to pump down deposition chambers when changing from one type of layer to another type of layer. The multi-chamber process minimizes the amount of time during which a completed reducing metal layer is exposed to oxygen, quickly transitioning from one deposition chamber to the next, resulting in the fuzzy interface layer. The primer provides at least the reliability of conventional primers while also taking advantage of the stability of thermite. The primer is useful not only for firearm cartridges, but also for other projectiles such as artillery, grenades, and other explosives and munitions. The primer is also useful for certain nail guns or other fastener guns which utilize primer-initiated propellants. One example of the primer will fit within a space designed for a conventional primer.
- (38) A variety of modifications to the above-described embodiments will be apparent to those skilled in the art from this disclosure. Thus, the invention may be embodied in other specific forms without departing from the spirit or essential attributes thereof. The particular embodiments disclosed are meant to be illustrative only and not limiting as to the scope of the invention. The appended claims, rather than to the foregoing specification, should be referenced to indicate the scope of the invention.

### **Claims**

- 1. A layered thermite composite, comprising: a substrate having a deposition surface and a rear surface; alternating layers of metal oxide and reducing metal deposited upon the substrate, the alternating layers of metal oxide and reducing metal being structured to react with each other in response to an impact applied to the rear surface of the substrate; and a fuzzy interface layer disposed between each metal oxide layer and reducing metal layer, the fuzzy interface layer containing reducing metal, reducing metal oxide, and metal oxide, the reducing metal, reducing metal oxide, and metal oxide being at least partially mixed together.
- 2. The layered thermite composition according to claim 1, wherein each of the reducing metal and metal oxide are present within each fuzzy interface layer in a gradient structure, the gradient structure being a majority reducing metal and reducing metal oxide adjacent to each reducing metal layer, the gradient structure being a majority metal oxide adjacent to each metal oxide layer.
- 3. The layered thermite composition according to claim 1, wherein the reducing metal adjacent to and within the fuzzy interface layer resists oxidation and resists growth after completion of deposition and prior to ignition of the layered thermite composite.

- 4. A primer, comprising a substrate having a deposition surface and a rear surface; alternating layers of metal oxide and reducing metal deposited upon the substrate, the alternating layers of metal oxide and reducing metal being structured to react with each other in response to an impact applied to the rear surface of the substrate; and a fuzzy interface layer disposed between each metal oxide layer and reducing metal layer, the fuzzy interface layer containing reducing metal, reducing metal oxide, and metal oxide, the reducing metal, reducing metal oxide, and metal oxide being at least partially mixed together.
- 5. The primer according to claim 4, wherein each of the reducing metal and metal oxide are present within each fuzzy interface layer in a gradient structure, the gradient structure being a majority reducing metal and reducing metal oxide adjacent to each reducing metal layer, the gradient structure being a majority metal oxide adjacent to each metal oxide layer.
- 6. The primer according to claim 4, wherein the reducing metal adjacent to and within the fuzzy interface layer resists oxidation and resists growth after completion of deposition and prior to ignition of the layered thermite composite.
- 7. The primer according to claim 4, further comprising at least one carbide-containing ceramic layer within the alternating layers of metal oxide and reducing metal, whereby, when the alternating layers of metal oxide and reducing metal react with each other, the at least one carbide-containing ceramic layer is ignited by the reaction between the reducing metal and metal oxide.
- 8. The primer according to claim 7, further comprising an adhesion layer separating each of the at least one carbide-containing ceramic layers and the layers of metal oxide or reducing metal which are adjacent to each of the at least one carbide-containing ceramic layers.
- 9. A cartridge for a firearm, the cartridge comprising: a casing having a front end, a back end, and a hollow interior; a bullet secured within the front end of the casing; a propellant disposed within the hollow interior; a primer secured within the back end of the casing, the primer being in communication with the propellant, the primer comprising; a substrate having a deposition surface and a rear surface; alternating layers of metal oxide and reducing metal deposited upon the substrate, the alternating layers of metal oxide and reducing metal being structured to react with each other in response to an impact applied to the rear surface of the substrate; and a fuzzy interface layer disposed between each metal oxide layer and reducing metal layer, the fuzzy interface layer containing reducing metal, reducing metal oxide, and metal oxide, the reducing metal, reducing metal oxide, and metal oxide, the reducing metal, reducing metal oxide, and metal oxide together.
- 10. The cartridge according to claim 9, wherein each of the reducing metal and metal oxide are present within each fuzzy interface layer in a gradient structure, the gradient structure being a majority reducing metal and reducing metal oxide adjacent to each reducing metal layer, the gradient structure being a majority metal oxide adjacent to each metal oxide layer.
- 11. The cartridge according to claim 9, wherein the reducing metal adjacent to and within the fuzzy interface layer resists oxidation and resists growth after completion of deposition and prior to ignition of the layered thermite composite.
- 12. The cartridge according to claim 9, further comprising at least one carbide-containing ceramic layer within the alternating layers of metal oxide and reducing metal, whereby, when the alternating layers of metal oxide and reducing metal react with each other, the at least one carbide-containing ceramic layer is ignited by the reaction between the reducing metal and metal oxide.
- 13. The cartridge according to claim 12, further comprising an adhesion layer separating each of the at least one carbide-containing ceramic layers and the layers of metal oxide or reducing metal which are adjacent to each of the at least one carbide-containing ceramic layers.