High-Temperature Solid Lubricants and Self-Lubricating Composites: A Critical Review

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Abstract: Solid lubricants are described as solid materials of intentionally introduced or in situ formed on contact surfaces in relative motion for the purpose of lowering friction and wear and providing protection from damage. Solid lubricants and advanced self-lubricating materials are widely used in modern industries, especially in aerospace, aviation, automotive, metallurgy, materials forming, and machining industries, and have attracted great interest in lubrication applications under very severe circumstances such as elevated temperatures, heavy loads, ultrahigh vacuum, extreme radiation, strong oxidation, and chemical reactivity environments. Many efforts have been made to develop self-lubricating composites by a variety of material preparation techniques, which include powder metallurgy, physical/chemical vapor depositions, thermal spraying, electrodeposition, laser cladding, and additive manufacturing. Although several reviews on the development of high-temperature solid lubricants have been published, most of them only focus on a type of material, a specific process, or application. In this paper, a comprehensive review is provided to present the state-of-the-art progress in solid lubricants, self-lubricating composites/coatings, and their effective functions that can be used over a wide variety of environmental conditions, especially at elevated temperatures. The solid lubricants considered include representative soft metals, layered structure materials (e.g., graphite, hexagonal boron nitride, transition metallic dichalcogenides, MAX phase), chemically stable fluorides, binary or ternary metallic oxides, especially alkaline earth chromates, and sulfates, and synergistic effects from these solid lubricants. This paper also provides new insights into design considerations of environmental adaptive solid lubrication, and the challenges and potential breakthroughs are further highlighted for high-temperature solid lubrication applications.

Keywords: solid lubricants; self-lubricating composites; friction; wear; extreme environments

1. Introduction

Solid lubricants are described as solid materials of intentionally introduced or in situ formed on contact surfaces in relative motion for the purpose of lowering friction and wear and providing protection from damage. Solid-lubricating materials for modern machinery, which uses rolling and sliding contact surfaces, mainly serve in extremely harsh conditions, such as elevated temperature, alternation of moist air atmosphere and vacuum, heavy load, high speed, strong oxidation and chemical reactivity, and severe thermal shock environments [1–5]. High friction, excessive wear, severe oxidation, and premature failure will inevitably occur if no lubrication mechanism is provided associated with the above-mentioned operation conditions. Nowadays, humanity is facing new challenges in the field of advanced solid lubrication, and novel self-lubricating components are required to operate durably and reliably in different industries such as aerospace, aviation, power generation, automotive, metallurgy, hot metal processing, and cutting tools.

Research in solid lubrication is understandably the minimization and elimination of materials and energy losses where sliding and rolling contact surfaces in relative motion are...
involved. Effective solid lubrication will bring greater materials and energy savings, higher efficiency, better comprehensive performance, and fewer machinery failures toward more reliable operations when exposed to extreme environments. Nowadays, there is a great need for lubricating polymeric, metallic, or ceramic components in sliding and rolling contacts, such as bearings and bushings for space satellites, gas turbine seals, bearings and variable stator vane bushings, and cylinder wall/piston ring lubrication for specific engine types from cryogenic temperatures to high operating temperatures [1,2,6–8]. For next-generation propulsion systems, a significant goal in the tribological design of advanced engines, bearings, and seals is to reduce undesirable friction and wear and attain a friction coefficient lower than 0.2 and a wear rate smaller than $10^{-6}$ mm$^3$/(Nm), which is expected to be independent of sliding velocity, applied load and ambient temperature for further increasing the efficiency of engines and reducing the emission of NO$_x$ and CO$_2$ [9].

However, developing new tribo-materials with a low friction coefficient and a small wear rate over a broad temperature range (i.e., room temperature to 1000 °C or even higher) has always been considered to be quite difficult and extremely challenging [10]. Higher operating temperatures in advanced power generating systems will provide improved efficiency, increased effectiveness, and reduced environmental impact where severe challenges are put forward to the existing solid lubricants, particularly from the point of view of high-temperature structural and chemical stability or surface lubricity intermittently but reliably [11]. Different motion mechanisms in nuclear power, hypersonic vehicles, and ballistic missiles also involve a very high operating temperature environment [12].

In the literature survey of solid lubricants and self-lubricating materials/coatings, the following classes of compounds are generally considered: polymer composites, soft metals, layered materials (graphite, hexagonal boron nitride, and transition metal dichalcogenides), alkaline earth fluorides, binary oxides (PbO, B$_2$O$_3$, Magneli phases TiO$_2$, V$_2$O$_5$, and MoO$_3$), and multi-component oxides (molybdates, tungstates, vanadates, tantalates, chromates, sulfates, silicates), oxythiomolybdates, MAX phases (Ti$_3$SiC$_2$, Ti$_2$AlC, etc.) [1,2,6–8,13,14].

High-temperature solid lubricants can be directly applied to the surfaces of machinery components by simple methods, such as painting and burnishing. A variety of fabrication methods of high-temperature self-lubricating materials or coatings include powder metallurgy processes such as pressureless sintering, hot pressing (HP), hot isostatic pressing (HIP), and spark plasma sintering (SPS), and coating processes such as electrodeposition, thermal/plasma spraying, magnetron sputtering, pulsed laser deposition, chemical vapor deposition, laser cladding, and additive manufacturing [2,8,15–18].

There have already been some reviews of various solid lubricants and preparation methods. However, few reviews deal with tribological design, diverse industrial applications of self-lubricating materials/coatings, and their underlying mechanisms, especially under extreme environmental applications [1–5]. The solid lubricants in this review concentrate on soft metals (Ag, Au), layered graphite and boron nitride (h-BN), transition metallic sulfides (MoS$_2$, WS$_2$), chemically stable fluorides (CaF$_2$, BaF$_2$), binary and multi-component oxides including silver-containing oxides, chromates and sulfates, and combinations of various solid lubricants. As it is difficult to investigate all compounds in these classes, this review proposes general design considerations relevant to environmental adaptive solid lubrication and provides a comprehensive understanding of self-lubricating materials and their effective function involved in challenging high-temperature environments.

2. Mechanisms of Solid Lubrication

In the case of no liquid/gas/grease lubrication, considerable adhesion exists between the rubbing surfaces of solid contact if two solid surfaces are clean (fresh metallic surface or in ultra-high vacuum) and all of the chemical films and adsorbates are removed. Generally, adhesion is regarded as either a physical or chemical interaction in nature [19]. In sliding and rotating machinery applications, adhesion usually results in friction and wear. Strong adhesion at tribo-stressing surfaces always leads to serious friction damage, cold welding, scuffing, or even breakdown, such as gears and bearings when subjected
to heavy loads, high velocities, and elevated temperatures. Adhesion depends mainly upon material pairs such as crystal structure, crystallographic orientation, mutual solubility, chemical activity, and separation of charges, as well as interface conditions such as surface cleanliness, normal load, temperature, atmosphere, duration of contact, and velocity [19]. In a hot metal forming process [4], components are subjected to extremely high thermal and mechanical dynamic loads and high velocity, which results in severe plastic deformation wear, and fatigue. Clearly, proper lubrication at elevated temperatures is needed to lower the friction stresses during forming and to avoid direct metallic contact, seizure, and galling. In a high-speed dry machining process, tool wear results from various wear mechanisms including adhesive, abrasive, chemical (by thermal diffusion), and electrochemical wear [4,5,19]. In the case of pantograph contact strips for electric railways [4], wear is derived from high-temperature mechanical impact, adhesion, and particle transfer due to arc discharge attack. For the thin-strip steel casting process, refractory side dams operate under severe environments of mechanical load, corrosion, wear, and thermal shock at elevated temperatures. Abradable seals used in the high-temperature compressor and turbine sections of gas turbine engines have to struggle against the degradation from adhesive, abrasive, thermal/corrosive, fatigue wear, and blistering. The failure modes of rolling-contact bearings, which contain substantial sliding for operating in a vacuum and high temperatures, are fatigue spalling under cyclic contact stressing and severe adhesive wear, commonly called scuffing or smearing [20].

Lamellar solids with weak interplanar cohesive bonds have a strong anisotropy of mechanical properties. Therefore, cleavage occurs generally at low shear stresses in some materials with anisotropy of mechanical properties and finally results in a distinctly reduced friction at the interface during sliding. These lamellae become self-lubricating due to the crystallographic slip mechanism at low shear stresses [21]. Similarly, conventional solid lubricants (such as MoS$_2$, graphite, etc.) have an easy-to-shear layered structure to provide self-lubricity; however, they generally become ineffective mainly due to oxidation-induced structural degradations in an oxidizing atmosphere (oxidation onset temperatures of 350 °C for MoS$_2$ and 450 °C for graphite) at elevated temperatures [11].

Non-lamellar soft solids, such as In, Pb, Sn, Ag, Au, and other soft compounds such as chemical stable fluorides, can lower friction and wear effectively when used as a strongly adherent thin film to the rubbing surfaces. Thus, the second solid lubrication mechanism is related to the formation of continuous and adherent soft solid films on a hard substrate during sliding. In this case, the shear strength of contacting asperities is determined by the softer thin solid film; however, the contact area is determined by the hard substrate. Clearly, the frictional force determined by theasperity shear strength and contact area becomes quite low under such conditions. The actual contact is between effective soft solid lubricants themselves due to the formation of a strongly adherent transfer film on the contact surface after a short running-in period. For example, a thin transfer film of lead is formed to provide self-lubricity for the lead-based alloys during sliding, while a low shear strength film is provided by the silver and fluoride eutectic constituents for the plasma-sprayed Ni$_8$Cr$_{20}$Cr$_2$O$_3$-Ag-CaF$_2$/BaF$_2$ composite coatings (PS304). They exhibit low coefficients of friction in dry sliding and are commonly used as bearing and seal materials [1,7,19].

The above-mentioned two fundamental modes of solid lubrication based on lamellar solids and soft films have been widely used for various sliding and rolling contact components. In addition, a variety of metals, intermetallic compounds, or ceramics react in air or water vapor atmospheres to some extent to form lubrious tribo-chemically reacted films. These tribo-chemically reacted films act as a low-shear-strength film and lead to low friction, which effectively separates the two metallic or ceramic surfaces from direct contact. Examples of vanadium or chromium as alloying elements in metals or nitride coatings form tenacious and lubrious oxide films, which are also responsible for low friction at elevated temperatures [2,22].
Although most oxides generally have poor lubrication properties due to their strong chemical bonds and brittle features, microstructurally designed thermally stable oxides may provide extreme high-temperature lubrication applications, especially in the oxidizing atmosphere. The lubrication mechanisms associated with lubricious oxides include the following seven aspects: (1) easy-shearing ability due to screening of cations by surrounding anions based on the crystal-chemical model \[10,23\]; (2) oxides are known to soften above the ductile-to-brittle transition temperature, which is typically 0.4–0.7 \(T_m\) where \(T_m\) is the melting point (in K). Material softening and plastic smearing contribute to the lubricious behavior observed for typical soft oxides once the operating temperature is attained to such a critical temperature \[24\]; (3) Low friction characteristic of viscous flow occurring from very thin liquid film, such as glass lubrication in hot metal forming process \[21,22,25,26\]; (4) creation and transfer of a reacted layer with lamellar structure and weak interplanar cohesive bonds (similar to the easy-shearing mechanism occurring in graphite and \(\text{MoS}_2\)) by oxidation or tribo-chemical reaction \[27\]; (5) intracrystalline slip deformation mechanism in textured nanocrystalline grains due to dislocation glide \[28,29\]; (6) The strain hardening at room temperature generally increase the friction stress during sliding, however, grain boundary sliding and grain rotation occurring in the microstructurally-refined surficial layer reduce the resistance to slide between the rubbing surfaces at elevated temperatures, and also contribute to the self-lubricity \[26\]; (7) In situ formation of glazed film with ultrafine nanograins by thermo-mechanically deformation-induced dynamic recrystallization and tribo-chemical reaction of surficial layer, which contributes to low friction at elevated temperatures \[26\]. Voevodin et al. \[30\] concluded that the primary adaptive lubrication mechanisms over a wide environmental range are associated with environmental-assisted oxidation to form easy-to-shear and low-melting-point binary \(\text{TiO}_2\), \(\text{V}_2\text{O}_5\), \(\text{MoO}_3\) Magneli phases and ternary oxides including silver molybdates, vanadates, tungstates, niobates and tantalates, temperature-activated diffusion or melting of soft metals, and thermo-mechanically-induced phase transitions to reorient hexagonal solids and promote surface self-hardening. Franz et al. \[31\] also summarized that the addition of V to nitride hard coatings enables a self-adaption mechanism via the formation of lubricious oxides at elevated temperatures, which improves the overall wear resistance and applicability of these coatings in metal cutting, in particular under dry-cutting conditions. A new approach is proposed to overcome brittleness in ceramic materials at relatively low temperatures \[32,33\]. Polycrystalline oxide lubricants with a reduced grain size of only a few nanometers may become more ductile during sliding and rolling contact. In this case, nano-sized oxides produce plastic deformation under tribo-stressing, in large part, due to grain boundary sliding or grain rotating at relatively low temperatures \[34\]. Thus, adaptive and lubricious oxide nanofilms introduced or generated on the rubbing surfaces may exhibit large plastic deformation and viscous flow to expand their lubricity over a broader operating temperature range.

3. Characteristics of Solid Lubricant Materials

As the primary function of solid lubricants and self-lubricating materials/coatings is to reduce friction and wear in sliding and rotating machinery, they must have the basic properties for effective lubrication and provide protection from damage over a wide variety of environmental conditions. In addition to superior lubricity, a suitable solid lubricant material should possess certain specialized properties such as high oxidation resistance, excellent thermal and chemical stability, high thermal conductivity, and low shear strength, particularly for specific engineering applications. Solid lubricant materials can be used in the form of bulks or coatings/films as well as dry powders, dispersions in lubricating oils or greases, and self-lubricating composites/coatings impregnated with solid lubricants.

The properties of an environmentally adaptive solid lubricant material include the following \[1,11,26\]: (1) Solid lubricants have low friction and moderately small wear rate in sliding or rolling contact without external lubrication from liquids or greases, and they must
possess low shear strength either by easy-to-shear lamellar microstructure such as graphite, MoS$_2$, and h-BN or by increased ductility at elevated temperatures such as Ag, Au, and CaF$_2$/BaF$_2$ eutectic. (2) Easy-to-shear alone does not ensure sufficient lubricity if the film does not strongly adhere to the contact surface. The solid films formed on the tribo-stressing surfaces must have a strong physical or chemical bond with the surface, which must remain continuous and intact during sliding or rotating processes such as physical vapor deposit MoS$_2$ films in non-oxidizing environments. (3) As surface heating caused by heavy loads or high sliding velocities results in the generation of easy-to-shear film and even localized melting, effective solid lubricants must remain thermally stable up to elevated temperatures such as thermally stable alkaline earth fluorides or sulfates. (4) The self-lubricating film must have good thermal conductivity to dissipate frictional heating from the contact region such as carbon-graphite parts impregnated with metallic fillers. (5) The self-lubricating film should function effectively between surfaces when subjected to high unit pressures and high velocities such as self-adaptive ultrahard multilayered or nanocomposite coatings. (6) Solid lubricants must be inert chemically to environments containing reactive vapors or fluids such as BN-based side-dam materials incorporated with $n$-ZrO$_2$ and SiC used for twin roll strip steel casting, and structural stability when subjected to radiation from radioactive sources such as high irradiation tolerant self-adaptive YSZ-doped MoS$_2$ nanocomposite films. (7) The self-lubricating film must have excellent oxidation/corrosion resistance at elevated temperatures such as plasma-sprayed Ag-CaF$_2$/BaF$_2$–containing composite coatings at elevated temperatures. (8) The self-lubricating film must have good electrical conductivity for the lubrication of sliding electrical contacts or brushes at elevated temperatures such as noble metal-, graphite- or MAX phase-based electrical contact materials. (9) Solid lubricants are able to provide safe operation, non-toxicity, and environmental compatibility, such as avoiding the usage of toxic PbO lubricant.

### 4. Classification of Solid Lubricants

Solid lubricants and self-lubricating solids are solid materials introduced or in situ generated between two rubbing surfaces, which exhibit low friction and wear in sliding or rotating machinery without external lubrication from liquids or greases. The most commonly used solid lubricants are polytetrafluoroethylene (PTFE), soft metals, graphite, and molybdenum disulfide [1–5,11,19]. Although these lubricants have been widely used singly or in various combinations, each of them has certain limitations. These limitations of common lubricating solids have stimulated the synthesis of novel lubricious compounds and composites/coatings with self-lubricating properties.

#### 4.1. Polytetrafluoroethylene (PTFE) and Polyimides

Many polymers have low densities, low friction coefficients, high chemical stability, and excellent machinability, and they are generally used under the conditions of cryogenic temperatures and vacuum [11]. Polytetrafluoroethylene (PTFE) and polyimides are the currently widely used solid lubricants in the family of polymers due to their good thermal stability and excellent tribological properties under different environmental conditions. However, these polymers exhibit poor thermal conductivity, high thermal expansion, and low radiation stability, and they are usually subjected to excessive cold flow under load (particularly at elevated temperatures), which leads to a lack of both strength and dimensional stability due to relatively poor heat dissipation efficiency [11]. PTFE, a fluorocarbon $(\text{C}_2\text{F}_4)_n$, is a crystalline polymer with a melting point of 325 °C. The simple zigzag backbone of −CF$_2$−CF$_2$− groups has a gentle twist of 180° over a distance of 13 CF$_2$ groups. The lateral packing of these rodlike molecules is hexagonal with a lattice constant of $a = 0.562$ nm [19]. The easy slippage of these rodlike molecules parallel to the c axis and their easy transfer onto the sliding partner account for the low friction characteristic of PTFE.

PTFE in the unfilled or filled forms retains good mechanical properties and lubricity up to 260 °C and can be used as a bearing material, which has the lowest coefficient of friction among any known solid lubricants, including graphite or MoS$_2$ [11]. A commonly adopted
value is 0.04 for the friction coefficient of PTFE against steel, while it can be as low as 0.016 at very high loads [11]. The PTFE coating on finger seals exhibits excellent wear resistance for reducing wear rates by 39%, which provides a valuable reference for the design of finger seals [35]. PTFE is usually reinforced with additives or filler reinforcements such as powdered graphite, MoS₂, or graphite fluoride (CF₃) in order to reduce the cold flow under severe load and speed conditions. Although these powdered additives can enhance tribological properties, they generally reduce their load-carrying capability. In this case, fiber reinforcement is used to meet the needs of maximum load capacity [36]. The fibers in the form of woven fabric are usually used to improve the creep resistance of bonded PTFE liners in airframe bearings and some heavily loaded bearings. From tests of nonmetallic plain cylindrical bearings, which were made of aramid (Kevlar) fiber-reinforced shell with a bonded, self-lubricating woven liner of polyester fiber-reinforced PTFE, low friction and wear were achieved up to 121 °C with a load capacity in excess of 207 MPa. However, the bearings are limited by creep deformation at higher temperatures [1].

Polyimide, used especially as coatings and films, are a class of imide-group-containing polymeric synthetic resins resistant to wear, corrosion, and high temperatures. The additions of CF₃ or MoS₂ solid lubricants to the polyimide varnish not only reduce the friction and wear of the coatings at room temperature but also eliminate completely the influence of molecular relaxation transition on tribological properties up to 500 °C. The polyimide-bonded CF₃ films are found to be excellent backup lubricants for foil gas bearings up to about 350 °C. Interestingly, polyimides are resistant to most common chemicals and solvents, but they easily suffer from the attack of alkalies.

Polyimides were widely used in bearings, bushings, gears, seals, and mechanical parts operating below 300 °C, at GE Aircraft Engine, Pratt & Whitney, Rolls & Royce, for example, the Vespel CP-8000 from DoPont™ for stator bushings of compressor in BR710 engine. Graphite fiber-reinforced polyimide (GFRPI) has attracted great attention for aviation applications and is being widely investigated for airframe and gas turbine engine applications due to the high strength, high thermal conductivity, and lubricity of graphite fibers. The Fibercomp, a chopped fiber/graphite-reinforced polyimide, from Foundry Service & Supplies, Inc., Ontario, CA, USA, has a friction coefficient of 0.1 to 0.2 and compressive strength of 172 MPa at 260 °C. The upper limit temperature for lubricious polyimides in the air is 350 °C, depending upon the type of polyimide. Polyimides are prone to becoming brittle and finally lead to wear damage by surface brittle fracture [11,37].

Interestingly, three-dimensional (3D) polyimide architectures with low volume shrinkage of only about 6% are fabricated by the UV-assisted direct ink writing (DIW) method, which provides the potential for three-dimensional printing of polyimides in the fields of aerospace, aviation, automobiles, and microelectronics [38]. The 3D target-region-lubrication printing of PTFE-filled photosensitive polyimide (PSPI) with excellent tribological properties for self-lubricating devices was realized by digital light processing and post-heat treatment. The PSPI-7wt.%PTFE composites fabricated by 3D printing exhibit a sharp reduction in friction coefficients by 88% and wear rates by 98%, respectively, and excellent mechanical properties with tensile strengths higher than 90 MPa, thermal stability up to 384 °C, and interlayer bonding as well [39]. For example, the friction coefficients are reduced to 0.09 for surface-lubricating and 0.04 for alternate-lubricating at 20 N. Importantly, a 3D-printed target-region-lubricating bearing was demonstrated effectively [39].

4.2. Soft Metals

There are some pure metals such as In, Sn, Pb, Zn, Ag, Au, and Pt, which are soft enough to be considered solid lubricants with their main lubrication mechanisms of enhanced ductility and the formation of an easy-to-shear tribolayer. The melting points and Mohs hardness values of these typical metals are shown in Table 1 [26]. In, Sn, Pb, and Zn have low melting points and Mohs hardness. Soft metal films of In, Sn, Pb, Zn, and their alloys with low melting points contain multi-slip systems and are able to compensate for microstructural defects effectively by frictional heat, which contributes to their
self-repairing ability as solid lubrication coatings under lightly loaded conditions and at relatively low temperatures. However, Ag, Au, and Pt have high melting points and low Mohs hardness [26]. Ag, Au, and Pt films with high melting points are easy-to-shear with a relatively inert nature to ambient vapor and temperatures, for a wide range of tribological applications such as long-term effective lubrication in X-ray tubes and satellites. Table 2 represents the fabrication methods and tribological behavior of self-lubricating materials containing soft metals [40–44].

Sn-Co binary alloy coatings are widely used for the substitution of hard chromium coatings in engineering applications. Ion-plated Pb coatings are especially designed for slow-rotating rolling element bearings in space mechanisms and have a higher cycle life than sputtered MoS$_2$ coatings in rolling contact bearing applications due to the unavoidable presence of lubricious PbO formed within the coatings. Pb-Sn-Cu in the form of plating on steel has been used as a lubricant for many years. The lubricious Pb-Sn-Ag and Ag-Cu-Pb-Sn additives were well distributed into the TiC-reinforced high-speed steel pre-forms to form an interpenetrating network microstructure for reducing friction and wear at elevated temperatures, and the wear rate can even be reduced by two orders of magnitude at 300–700 °C [45].

Table 1. Melting points and Mohs hardness values of typical soft metals [26].

| Material | Melting Point | Mohs Hardness |
|----------|---------------|---------------|
| In       | 155 °C        | 1.0           |
| Sn       | 232 °C        | 1.8           |
| Pb       | 328 °C        | 1.5           |
| Zn       | 419 °C        | 2.5           |
| Ag       | 961 °C        | 2.5           |
| Au       | 1063 °C       | 2.5           |
| Pt       | 1755 °C       | 4.3           |

Table 2. Fabrication methods and tribological behavior of self-lubricating materials containing soft metals [40–44].

| Materials       | Fabrication Method       | Tested Conditions                      | Results/Observations                                                                 |
|-----------------|--------------------------|----------------------------------------|--------------------------------------------------------------------------------------|
| TiN-In [40]     | Sputtering deposition    | Pin-on-disk; Al$_2$O$_3$ ball; load 1 N; 0.1 m/s; 150–1200 °C | • $\mu$: 0.5–0.6                                                               |
|                 |                          |                                        | • TiN-In coating has high sustainability of lower friction coefficient for more than 1500 cycles. |
|                 |                          |                                        | • TiN-In coating exhibited four times longer wear life than TiN coating in wet machining. |
|                 |                          |                                        | • Friction coefficient significantly increased due to the indium oxidization above 450 °C. |
|                 |                          |                                        | • $\mu$: ~0.3 (20–600 °C), 0.09 (800 °C)                                             |
|                 |                          |                                        | • $W$: $1.4 \times 10^{-5}$, $7.7 \times 10^{-4}$, $6.0 \times 10^{-5}$ mm$^3$/(Nm) (20, 600, 800 °C) |
| NiMoAl-Ag [41]  | High-velocity oxy-fuel spraying | Ball-on-disk; Si$_3$N$_4$ ball; load 5 N; 0.1 m/s; 20–800 °C | • Enriched silver on sliding surface form a lubrication film below 400 °C. |
|                 |                          |                                        | • A synergistic effect of molten Ag and Ag$_2$MoO$_4$ by tribochemical reactions gives a high lubricating property at high temperature. |
Table 2. Cont.

| Materials                  | Fabrication Method                        | Tested Conditions                                                                 | Results/Observations                                                                 |
|----------------------------|--------------------------------------------|-----------------------------------------------------------------------------------|--------------------------------------------------------------------------------------|
| NiMoAl-Al$_2$O$_3$-Ag [42] | Plasma spraying                            | Ball-on-disk;                                                                    | • $\mu$: 0.53, 0.36, 0.17 (RT, 500, 900 $^\circ$C)                                    |
|                            |                                             | Al$_2$O$_3$ ball; load 12 N;                                                      | • W: $1.47 \times 10^{-5}, 8.84 \times 10^{-5}$, $3.35 \times 10^{-5}$ mm$^3$/Nm (RT, 500, 900 $^\circ$C) |
|                            |                                             | 0.1 m/s; RT-900 $^\circ$C                                                          | • Continuous lubricating film mainly consisted of NiO and Ag$_2$MoO$_4$ formed at 900 $^\circ$C. |
|                            |                                             |                                                                                  | • Continuous lubricating film forming at high temperature improved subsequent friction performance during multitemperature cyclic tests. |
| Al$_2$O$_3$-DLC-Au-MoS$_2$ [43]| Magnetron-assisted pulsed laser deposition| Ball-on-disk;                                                                    | • $\mu$: 0.13–0.14 (air at 40% RH), 0.02–0.03 (N$_2$ at <1% RH), 0.1 (air at 500 $^\circ$C) |
|                            |                                             | M50 steel ball (RT); Si$_3$N$_4$ ball (500 $^\circ$C); load 100 g; 0.2 m/s;       | • Cycles to failure 500 $^\circ$C: >10,000                                          |
|                            |                                             | air at 40% RH and N$_2$ at <1% RH                                                 | • The higher ratio of hard phase contributes to extend coating wear life at 500 $^\circ$C. |
|                            |                                             |                                                                                  | • $\mu$: 0.2 (6% textured dimple density, between 25 and 600 $^\circ$C)             |
| Ta-Ag [44]                 | Magnetron sputtering deposition             | Ball-on-disk;                                                                    | • W: $5.2 \times 10^{-5}$ mm$^3$/Nm (at 600 $^\circ$C)                             |
|                            |                                             | Si$_3$N$_4$ ball; load 2 N; 0.128 m/s; 25–600 $^\circ$C                            | • Ta$_2$O$_5$ and silver lead to low friction coefficient at high temperature.       |
|                            |                                             |                                                                                  | • Textured dimples store lubricants to extend the wear life.                       |

Electrodeposition and physical vapor deposition processes have been applied to fabricate soft metallic coatings such as Ag, Au, and Pt, which are particularly valuable at very high temperatures and under severe conditions, such as in spacecraft. Ag and Au lubricating films deposited by oxygen-ion assisted screen cage ion plating can avoid undesirable subsurface cracking and reduce friction and severe wear. Herein, achieving low and steady friction coefficients and low wear is mainly attributed to the strong adhesion of lubricious Ag and Au films to alumina [46]. Self-lubricating Au and Ag films can be used in space capsules, advanced jet engines, and high-speed lightly loaded machines. An Au-Co alloy coating was designed for effectively lubricating ring tracks and rolling flexures in the roll ring assembly on space station freedom.

Composite brushes used in the solid lubricated slip ring assemblies on spacecraft consist mainly of a conductive metal such as Ag, Au, and/or Cu, incorporated MoS$_2$ and/or a small amount of graphite, which contributes to a balance of low friction and wear and low electrical contact resistance. The graphite was originally added into the conductive metallic matrix to generate self-lubricity for military aircraft, where both MoS$_2$ lubrication in low humidity at high altitude and graphite lubrication in humid air at low altitude are needed. Clearly, incorporating multiple solid lubricants such as soft metals, graphite, and MoS$_2$ can achieve reversible surface chemistry and morphology during sliding that is well-suited for lubrication over a broad range of ambient environments.
Au and Ag lubricants can be incorporated into high-temperature intermetallic or ceramic matrix composites/coatings for providing lubrication from room temperature to 600 °C [3]. They have high diffusion coefficients, which help the easy formation of lubrication films onto the surfaces of intermetallic or ceramic matrix composites, which are used to lubricate bearings, seals, and fasteners in sliding or rolling contact, and exhibit good thermochemical stability up to elevated temperatures [5]. Figure 1 shows the worn surfaces of the cermets infiltrated with and without Ag lubricant. The Ag-infiltrated cermet exhibited a thin lubricating film on the surface; however, obvious furrow marks were observed on the unmodified cermet. Friction heat and surrounding temperature induce the formation of a melting film, which can effectively reduce the friction coefficient [47]. Typical examples of Au- and Ag-containing coatings for tribological applications under the harshest conditions are YSZ/Au/MoS$_2$/DLC, YSZ/Ag/Mo, and Al$_2$O$_3$/DLC/Au/MoS$_2$ nanocomposite films fabricated by hybrid magnetron sputtering and pulsed laser deposition processes. All these films have friction coefficients of 0.1–0.4 for from 5000 to 10,000 cycles at 500 °C due to the migration of easy-to-shear noble metals to the surface to provide a lubricious layer [3,43,48,49].

![Figure 1](image_url)

Figure 1. Morphologies of worn surface of the cermets infiltrated with and without Ag lubricant at $p = 50$ N and $T = 800$ °C. (a) without Ag lubricant added; (b) Ag-infiltrated. Reproduced with permission from Reference [47], Copyright © Elsevier B.V. 2006.

Since the lubrication films of soft metals are easily removed by shear, they generally have poor wear resistance and a short useful lifetime under severe tribo-stressing, particularly after becoming softened at elevated temperatures. This challenge can be partially overcome by intentionally designing the reservoirs of soft metals in hard textured surfaces and gradually releasing these softer materials during sliding to generate an easy-to-shear transfer film when needed [50]. Examples are the incorporation of silver [44] or in combination with MoS$_2$ [51] as solid lubricants applied to a textured hard surface to reduce friction and wear.

In addition, some metals such as Fe, Cu, Ni, Mo, and Cr exhibit high coefficients of friction at room temperatures, but considerable improvements in friction reduction to only half of their original values are noted during sliding above their oxidational temperatures [26]. This improvement in friction during sliding is associated with the formation of oxidized products in the wear track for these metals. Therefore, these soft oxides formed on the surfaces of metals and alloys provide the most effective friction-reducing characteristics and separate the rubbing surfaces from direct metallic contact. In recent years, pulsed laser deposition, plasma spraying, magnetron sputtering, and ion bombardment assisted deposition (IBAD) have been employed to produce Au/Cr, Zn/W, Ni/Ti, Cu/Mo, W/MoS$_2$, and AlCuFeCr multilayered films for tribological applications, such as turbomachinery components of bearings, seals, and fretting interfaces up to 580 °C.
4.3. Layer Lattice Solid Lubricants

4.3.1. Graphite

Graphite is hexagonal in orientation and is composed of planes of polycyclic carbon atoms. The bonding between the basal planes is weaker due to the presence of a longer distance between carbon atoms along the c axis, and the parallel layers of hexagons stack 0.34 nm apart. Close examination of a metallic mating surface in sliding against graphite reveals coefficients of friction ranging from 0.05 to 0.15 in an ambient environment. However, the coefficient of friction when sliding on a face perpendicular to the basal planes compared to sliding parallel to the basal planes is three times or even higher [19]. Therefore, if tested parallel to the basal planes, graphite is soft and lubricates in normal air; however, it fails to lubricate in a vacuum or at high altitudes and wears rapidly. In fact, its coefficient of friction in vacuum or dry nitrogen is typically ten times greater than in air. Both natural and synthetic graphite must adsorb moisture or other condensable vapor such as hydrocarbons in order to be slippery and lubricious. The bonding energy between the hexagonal planes of graphite is reduced to a lower level by the adsorption of water under high-humidity environments. In fact, graphite exhibits excellent lubrication in boundary lubrication conditions due to its good affinity for hydrocarbon lubricants. In an oxidizing atmosphere, graphite lubricates effectively up to 450 °C and then fails due to structural degradation by oxidation [26]. Powdered graphite lubricants entrained in a gaseous carrier were used for rolling-contact bearings and high-speed spur gears for operation at temperatures over 650 °C in military aircraft. Impregnated graphite parts are not suited for applications in which mechanical shock or loading is relatively high and are often used in high-temperature structural applications such as missile-nozzle inserts, heat shields, and fuel-chamber liners. Under some circumstances, graphite additives are able to provide very short-term lubrication up to 1000 °C or even 1200 °C in the hot metal forming process. Graphite is widely used as the rubbing element in mechanical seals and as electrically conducting brushes for motors and generators. Generators used in airplanes flying at high altitudes worsen the performance of graphite brushes due to the lack of moisture in the air and water vapor. In this case, the brushes wear out rapidly at high altitudes, and thus, oxygen and water vapor are considered to be the most important gases for the brushes.

For applications where only a minor lubricity is needed and a more thermally insulating coating is required, amorphous carbon is predominant. In this case, amorphous carbon and graphite can be combined to take full advantage of the strengths and weaknesses of each. Parts made from carbon-graphite are strong and hard and exhibit low friction. Some carbon-graphite bearings are capable of extended service at temperatures above 580 °C, which is ideal for chemically aggressive applications. Carbon-bonded graphite with high thermal conductivity is desirable for high-temperature applications with high sliding speeds up to 150 m/s (e.g., encountered in aircraft jet engine seals) in order to dissipate the frictional heat. Many carbon-graphite parts impregnated with fillers such as various metals (e.g., Ag, Cu-Pb alloys) and high-temperature chemical salts provide lower permeability, higher strength and hardness, low friction, wear and oxidation resistance in dry air at temperatures up to 750 °C. In a hot metal forming process, a high-temperature graphite-based solid lubricant consisting of amorphous silica, aluminum dihydrogen phosphate, and graphite exhibits strong adhesion to the substrate and excellent tribological properties with a low friction coefficient of about 0.05 and good wear protection for the workpiece at elevated temperatures in the air [52].

Graphite fluoride (CF$_x$)$_n$, also referred to as carbon monofluoride (when $x = 1$), is a solid lubricant that can be described as a layer lattice intercalation compound of graphite [1]. It is an electrical insulator, unlike graphite, and is non-wettable by water. Graphite fluoride is generally synthesized by the direct reaction of graphite with fluorine gas at controlled pressure and temperature. However, the frictional properties of graphite fluoride are less influenced by humidity than either MoS$_2$ or graphite. Graphite fluoride synthesized by the above-mentioned method exhibits grey to pure white depending upon
its composition parameter, where $x$ can vary from about 0.3 to 1.1. The fluorine to carbon bonds are covalent with the fluorine atoms located between the distorted basal planes. The spacing between the basal planes is expanded from 0.34 nm in graphite to 0.75 ± 0.15 nm in (CF$_2$)$_n$, which leads to a further decrease in shear and cleavage strength parallel to the basal planes [11]. (CF$_2$)$_n$ is not known to oxidize in air, but it decomposes thermally above about 450 °C to form carbon tetrafluoromethane, other low molecular weight fluorocarbons, and carbon. (CF$_2$)$_n$ exhibits extreme plasticity within lubricated contact.

### 4.3.2. Hexagonal Boron Nitride (h-BN)

h-BN, either in its pure form or as a composite, is an extremely suitable material for a variety of high-temperature applications, such as solid lubrication material, gas seals for oxygen sensors, high-temperature furnace parts, crucibles for molten glass and metals, as well as evaporation boats for aluminum and side dams for thin-strip casting [53]. h-BN, referred to as “white graphite”, is composed of a layered structure containing a network of (BN)$_3$ rings. It has very anisotropic shear properties with preferred shear parallel to the basal planes or perpendicular to the c-axis [11]. Lamellar slip along the basal plane is regarded as the prominent lubrication mechanism at high temperatures. However, h-BN has significant drawbacks such as weak adhesion to most metals and ceramics and the ability of difficult-to-sinter, which generally leads to low strength and low quality of composite materials [1,11]. The h-BN layers show a down-graded tribological performance due to relatively stronger van der Waals interlayer forces than graphite or MoS$_2$. However, it works better under humid and high-temperature conditions due to its stronger oxidation resistance and higher thermal stability, making it suitable for sintering processes. Especially h-BN is currently used as an additive to improve the tribological performance of ceramics and composites [54]. h-BN exhibits friction coefficients of 0.2-0.25 in a normal atmosphere, below 0.1 in humid air, and even smaller in water and Vaseline [55–57]. In order to improve the lubricity under different environmental conditions, h-BN as lubricating micro-particles are also impregnated into a porous surface [58,59] or added to lubricating oil [60] or even water [61]. The thermal stability of h-BN is better than that of MoS$_2$ or graphite, but its friction coefficient at room temperature is relatively high; however, it drops to about 0.15 at 600 °C. Interestingly, h-BN exhibits a good lubrication feature even at a service temperature of 1200 °C in an oxidizing atmosphere. Friction and wear tests of pure h-BN and h-BN-10 wt.% CaB$_2$O$_4$ was performed in sliding against Si$_3$N$_4$ counterpart on a ball-on-disk tribometer in atmospheric and water vapor environments from room temperature to 800 °C [56]. Both pure h-BN and h-BN-10 wt.% CaB$_2$O$_4$ have a similar friction coefficient of less than 0.2 in atmospheric conditions at room temperature. At 400 °C, both of them exhibit high friction coefficients of 0.58 and 0.51 due to the adhesion of h-BN on the coupled Si$_3$N$_4$ ball, respectively. On further increasing the test temperature to 800 °C, the coefficients of friction for pure h-BN and h-BN-10 wt.% CaB$_2$O$_4$ decreased to 0.38 and 0.35, respectively, which is attributed to the formation of molten B$_2$O$_3$. Interestingly, the friction and wear tests performed in the water vapor environment showed prominently reduced friction coefficients of 0.08 and 0.07 for pure h-BN and h-BN-10 wt.% CaB$_2$O$_4$ at room temperature, respectively. However, increasing the test temperature to 400 °C leads to an increase of the friction coefficient to between 0.25 and 0.23, which is distinctly lower than those tested under atmospheric conditions, due to the reaction of h-BN with water vapor to form B$_2$O$_3$ and finally H$_3$BO$_3$. The formed H$_3$BO$_3$ has a lamellar structure that can shear very easily under tribo-stressing to reduce friction and wear. At 800 °C, the friction coefficients for pure h-BN and h-BN-10 wt.% CaB$_2$O$_4$ are 0.22 and 0.21, respectively, which are also lower than those obtained under atmospheric conditions. Table 3 summarizes the fabrication methods and tribological behavior of self-lubricating materials containing h-BN [56,59,62–65].
Table 3. Fabrication methods and tribological behavior of self-lubricating materials containing h-BN [56,59,62–65].

| Materials                  | Fabrication Method | Tested Conditions | Results/Observations                                                                 |
|----------------------------|--------------------|-------------------|--------------------------------------------------------------------------------------|
| h-BN                       | Hot pressing       | Ball-on-disk;     | • μ: 0.18, 0.58, 0.38 (h-BN, atmosphere ambience, at RT, 400, 800 °C)                |
| h-BN-10 wt.% CaB₂O₄ [56]   | Hot pressing       | Si₃N₄ ball; load 1.5 N; 0.188 m/s; RT-800 °C | • μ: 0.08, 0.25, 0.22 (h-BN, water vapor ambience, at RT, 400, 800 °C)               |
|                            |                    |                   | • CaB₂O₄ as the sintering additive promoted crystallization of h-BN and decreased the friction coefficient. |
|                            |                    |                   | • H₃BO₃ formed by the reaction of water vapor with h-BN has a lamellar structure to reduce friction and wear. |
| Cu-based composites        | Hot pressing       | Block-on-ring;    | • μ: ~0.5                                                                             |
| (Cu-Sn-Al-Fe-h-BN-        |                    | AISI52100 bearing steel; load 50–125 N; 1.04–2.6 m/s; RT | • W: 1.3 × 10⁻⁵–4.3 × 10⁻⁵ mm³/(Nm) (Varying with the content of h-BN and graphite)    |
| Graphite-SiC) [59]         |                    |                   | • The higher ratio of graphite to h-BN contributed to form relatively continuous and compact tribo-films. |
| B₄C-h-BN [62]              | Hot pressing       | Pin-on-disk;      | • μ: 0.591–0.321 (content of h-BN rising from 0 to 30 wt.%)                          |
|                            |                    | B₄C pin; load 10 N; 0.656 m/s; 25 °C | • W: 2.07 × 10⁻⁵–1.94 × 10⁻⁴ mm³/(Nm) (content of h-BN from 0 to 30 wt.%)             |
| NiCr/Cr₂C₂-NiCr/           | Plasma spraying    | Ball-on-disk;     | • The transfer film formed on the wearing surface is considered as the main cause of tribological properties improvement. |
| h-BN [63]                  |                    | Si₃N₄ ball; load 9.8 N; 0.188 m/s; 20–800 °C | • μ: 0.65–0.55 (20–800 °C)                                                         |
| Ni-P-h-BN [64]             | Electroless plating| Pin-on-disk;      | • W: 5.3 × 10⁻⁵–1.15 × 10⁻⁴ mm³/(Nm) (20–800 °C)                                   |
|                            |                    | AISI52100 steel ball; load 2 N; 0.1 m/s; RT | • h-BN reduces the friction coefficient but weaken the wear resistance due to the poor cohesive strength. |
| NiCrWMoAlTi-i-h-BN-Ag [65] | Hot pressing       | AISI52100 steel ball; load 20 N; 1 m/s; RT-600 °C | • μ: 0.2                                                                            |
|                            |                    |                   | • W: 1.24 × 10⁻⁶ mm³/(Nm) Stable sliding distance > 1000 m                            |
|                            |                    |                   | • μ: 0.54–0.37 (RT-600 °C) Stable sliding distance > 1000 m                           |
|                            |                    |                   | • W: 7 × 10⁻⁵, 7 × 10⁻⁴, 2 × 10⁻⁴ mm³/(Nm) (RT, 200, 600 °C)                          |
|                            |                    |                   | • Stable sliding distance > 600 m                                                    |

The addition of BN or TiN into the Si₃N₄ matrix can reduce the friction and wear in the sliding of self-mated couples below 100 °C, especially at room temperature in humid air, where the friction coefficient is reduced to 0.1 and the wear rate is one-fifth that of monolithic Si₃N₄. Besides the optimized mechanical properties of Si₃N₄-BN and Si₃N₄-TiN composites without any glass phase, the distinct improvement in tribological behavior was attributed to the tribo-chemical reaction to form relatively soft lubricious oxide layers such as H₃BO₃ and TiO₂₋ₓ or an intrinsic solid lubricant of BN-H₂O [9,66]. For Si₃N₄-based cutting tools, chipping and subsequent fracture lead to unavoidable failure when subjected to fatigue loading and high temperatures, particularly during high-speed machining of hard-to-cut materials [67]. However, when h-BN is added to Si₃N₄, a significant tribological improvement is achieved with austenitic stainless steel as tribo-pair due to the formation of a tribo-film comprising Fe₂O₃, SiO₂, and B₂O₃. h-BN is used as a high-temperature solid
lubricant in the ceramic matrix such as Al₂O₃, TiB₂, and B₄C [62] due to its high chemical stability and resistance to oxidation. However, one challenge of using h-BN as the additive in the cutting tool is the difficulty to sinter of fully densification and poor adhesion with ceramics due to its flaky structure and low diffusion coefficient [68].

h-BN, as a clean solid lubricant, is able to replace dirty graphite or MoS₂ in Al-forming processes without staining. The tribological performance, including the lubrication-film stability and the surface quality, depends mainly upon the particle size and concentration of h-BN powders [69]. h-BN is chemically inert and difficult to wet with most molten metals. Therefore, h-BN is an important solid lubricant for various industrial applications, especially in aircraft turbo engines [56] and in metalworking processes where high-temperature lubrication and/or environmental cleanliness are preferentially required [60].

Layered-structure h-BN is incorporated with monoclinic zirconia or ZrO₂-SiC or SiAlON as a composite material for side dams in twin roll strip steel casting due to its unique combination of properties such as excellent release and lubrication properties, high-temperature thermal shock resistance, chemical inertness, and easy machinability to produce complex shapes from the as-sintered billets [53,70,71]. One of the main products of ESK ceramics GmbH & Co. KG, a subsidiary of Ceradyne Inc., is hexagonal boron nitride, which can be used as powders, coatings, or in its sintered form either as pure BN or as a composite at a maximum operating temperature of up to 1100/1500 °C in an oxidizing/inert atmosphere, respectively [53].

BN-based materials incorporated with monoclinic ZrO₂ and SiC are used in steel production as the best side-dam material for twin roll strip steel casting as well as a broken ring for horizontal continuous casting [53,72–74]. Wear tests were carried out from room temperature to 400 °C using a pin-on-disc high-temperature tribometer with BN-based composite pins sliding against a rotating disc of nickel [70]. Control of the wear process requires a better understanding of the mechanisms of third body formation and the velocity accommodation mechanisms. The wear surface of BN-based composites shows textured grooves, brittle fracture, and microplastic deformation [53].

C/C-h-BN-SiC composites were prepared by molding, carbonization, and liquid silicon infiltration processes, which exhibited excellent oxidation resistance, self-healing function, and self-lubrication properties for high-temperature applications. Herein, h-BN addition promotes the formation of a lubricious tribofilm and distinctly reduces the friction and wear of C/C-h-BN-SiC composites without any clamping stagnation phenomenon in high braking speed conditions [75].

h-BN is used as a solid lubricant or release agent either as a sintered body (e.g., side dams) or applied as suspensions or powders or coatings (e.g., aluminum extrusion or titanium shaping, aluminum casting, superior lubricity). A 15 μm thick h-BN coating prepared from a polyborazylene polymeric precursor was deposited on titanium alloys and annealed via infra-red irradiation in a rapid thermal annealing furnace for high-temperature tribological applications. The friction coefficient tested at 360 °C was reduced from 0.72 for Ti-alloys to 0.35 for the Ti@h-BN coating when sliding against the coupled 15-5PH stainless steel cylinders using a cylinder/disk configuration [76,77]. The Ni-P-35 vol.% h-BN autocatalytic composite coating has a wear rate in the order of 10⁻⁶ mm³/(Nm) and a friction coefficient of 0.2 against the tribopair of AISI52100 steel ball at ambient temperature [64]. The addition of silver and h-BN nanopowders into nickel-based composites exhibits a self-lubricity from room temperature to 600 °C [75]. The mechanical strength and tribological behavior of ion-beam-deposited h-BN films on non-metallic substrates were also investigated with diamond pin sliding experiments. BN films on Si and SiO₂ exhibited low friction coefficients of less than 0.1; however, BN films on nonmetallic substrates could fracture at a critical load [78].
4.3.3. Transition Metal Dichalcogenides

Transition metal dichalcogenides (TMD) solid lubricants include a wide range of hexagonal layered compounds formed by the bonding between chalcogenides such as sulfur, selenium, and tellurium, as well as transition metals such as molybdenum, tungsten, and niobium [79,80]. The lubrication mechanism for this class of solid lubricants is closely related to their microstructure, and the weak van der Waals adhesion forces between sulfur-like atoms lead to the formation of easy-to-shear lamellas [54]. Some compounds, such as Mo- and W-disulfides and Nb-diselenide, have a closely packed lamellar structure with strong bonding between the transition metal and chalcogenide ligands. MoS$_2$ and WS$_2$ are typical representatives of intrinsic solid lubricants and are the best lubricants of choice, especially for vacuum applications. They do not require adsorbed materials or additives to develop lubricating capability. However, when subjected to the adsorption of chemical compounds like H$_2$O from the environment, the lubricity worsens rapidly, which significantly influences the ability of the lamellas to slide against each other. It must be noted that a layered microstructure is not sufficient to achieve self-lubrication, and the strength of the interlayer bonding is a decisive factor, as observed for similarly layered TMDs like NbS$_2$, TiS$_2$, etc. One of the main prerequisites to achieving ultralow friction for MoS$_2$ lubricants is the absence of contaminants, such as oxygen, water, and hydrocarbons [81]. The susceptibility of MoS$_2$ films to moisture is reduced by adding different dopants, including Pb, Au, and polytetrafluoroethylene (PTFE). It is noted that the addition of carbon to burnished and bonded MoS$_2$ improves the performance of MoS$_2$ in humid air [3].

MoS$_2$ is thermally stable in non-oxidizing environments up to 1100 °C, but in the air, the onset oxidation temperature of MoS$_2$ is at around 350 °C. The oxidized product MoO$_3$ is believed to be abrasive, which limits the maximum-use temperature, although now it may be virtually innocuous. Friction coefficients for sputtered MoS$_2$ in a vacuum were reported to be as low as 0.01, however, oxygen and especially water vapor in the ambient atmosphere causes a slow oxidative degradation of MoS$_2$, which leads to early failure. The oxidation rate also depends strongly on the airflow rate through the reaction chamber. MoS$_2$ was oxidized to molybdic oxide (MoO$_3$) in 1 h at 400 °C at a low flow rate of $5.5 \times 10^{-6}$ mm$^3$/s and lost the lubricating ability in the air. Therefore, MoS$_2$ is a poor lubricant when used in humid air, and it is also corrosive to the mating metallic surface due to the formation of a surface layer comprising oxysulfide (MoOS$_2$) and sulfuric acid of H$_2$SO$_4$. WS$_2$ is a better lubricant than MoS$_2$ at high temperatures in both air and non-oxidizing atmospheres. Similarly, WS$_2$ starts to oxidize appreciably above 425 °C, which also limits distinctly its usefulness in air. The XPS analysis on the wear scar of WS$_2$ coatings in Figure 2 shows a significant difference in elemental distribution and oxidation state at temperatures of 100 °C and 500 °C. A very low W$^{6+}$ content from WO$_3$ peaks indicates slight oxidation of the surface on the wear scar at 100 °C. However, sulfur almost disappears at 500 °C, and WS$_2$ is completely oxidized to form WO$_3$ [82]. WS$_2$ nanoparticles with a closed-cage structure have already been investigated for superior lubrication applications under harsh circumstances, which provide excellent performance for the tribological contact between silicon nitride ball and alumina ceramic block due to gradual exfoliation of the fullerene-like WS$_2$ onions followed by transfer of monomolecular WS$_2$ sheets to the coupled surface [83].
TMD compounds can be used as lubricants themselves or as lubricant additives to reduce both friction and wear. TMD powders are frequently used in such applications as metal-forming dies, threaded parts, sleeve bearings, and electrical contacts in relays and switches. The applications of TMD-containing materials include bulk composites sintered by powder metallurgy, thin films deposited by PVD or burnishing, and thick coatings either by laser cladding or thermal spraying [54]. No matter what techniques are adopted for material fabrication, superior low friction and small wear rate must be provided in a wide operating temperature range to achieve the desired lubricity [2]. Refractory metal-based composites containing MoS$_2$ or WS$_2$ can be used at loads up to 70 GPa at 500 °C and 7 GPa at 800 °C in a vacuum, such as applications for bushings, seals, gears, clutch facings, and electric motor brushes. Applying MoS$_2$ as a solid lubricant onto solid surfaces includes burnishing, spray bonding, physical vapor deposition, and, in particular, sputtering, such as for applications of bearings employed in spacecraft, which contributes to maintaining essential lubrication properties, thus reducing friction and/or wear as well as enhancing their service life and energy efficiency by avoiding premature failures.

In the case of burnished MoS$_2$ or WS$_2$ coatings, the preferential orientation of MoS$_2$ or WS$_2$ crystallites can be introduced within the coating where the basal planes are mostly parallel to the sliding direction to achieve low friction, which is commonly used in bearings and other sliding/rolling applications. Bonded coatings can be applied by spraying to produce a more uniform coating with careful process control, such as the superior lubrication of gears for the space station remote manipulator system (SSRMS) without apparent wear over several million cycles. Solid lubrication techniques of MoS$_2$ films have been successfully adopted in representative ball bearings, pointing mechanisms, slip rings, gears, and release mechanisms in space applications [84]. The increasing solid lubrication of MoS$_2$ needs to operate in a wide range of temperature conditions for longer durations in a robust and reliable fashion for planned space missions, such as the ESA’s BepiColombo mission to Mercury, where long-term exposure to temperatures on the order of 250 °C is involved [85]. Herein, a particularly important example associated with the failure of the MoS$_2$-based solid lubricant during space operation is the catastrophic high-gain antenna.
deployment failure associated with the Galileo spacecraft [86], although the reliability of lubrication provided by MoS\(_2\) was successfully tested before deployment on Earth under ambient conditions. MoS\(_2\) was also determined as the solid lubricant of choice for precision instruments on the Hubble Space Telescope and its successor of the James Webb Space Telescope (JWST) [80].

Sputtered and resin-bonded MoS\(_2\) coatings are used in numerous applications of satellites and the space shuttle. Resin-bonded spray coatings followed by the heat-curing process are typically from 5 to 15 \(\mu\)m thick with friction coefficients of only from 0.06 to 0.15 and superior wear life, depending on humidity and sliding conditions. However, the main inorganic binders in inorganic-bonded MoS\(_2\) coatings used for space applications are generally silicates (e.g., \(\text{Na}_2\text{SiO}_3\)) and phosphates (e.g., \(\text{AlPO}_4\)), which can tolerate moderately elevated temperatures up to 650–750 \(^\circ\)C for various types of gears and low-cycle bearings on launch vehicles. However, one challenge is associated with softening or degradation in the presence of water/humidity. A phosphate-bonded MoS\(_2\) coating is used for the lubrication of the Mars Science Laboratory (MSL) Curiosity Rover Main Differential Pivot to deliver the frictional torque. Sputter-deposited MoS\(_2\) coatings incorporated with or without soft metals and other compounds exhibit considerably low friction for the lubrication of components in space programs. However, a main technical obstacle associated with these sputtered coatings is the difficulty of deposition on large parts in a vacuum [87].

4.3.4. Layered MAX Phase Materials

MAX phase materials include ternary MAX phases (\(\text{M}_{n+1}\text{AX}_n\)), MAX phase-like compounds (\(\text{M}_{n+2}\text{AX}_n\) and \(\text{M}_m\text{AX}_p\), \(m \neq n + p\)), quaternary MAX phases (\(\text{M}_n\text{AX}_p\) and \(\text{M}_m\text{AX}_p\)) and high-entropy MAX phases such as \(\text{M}_2\text{AX}\) (\(\text{M} = \text{Ti}, \text{Zr}, \text{Hf}, \text{Nb}, \text{Ta}\)) or \(\text{V}_2\text{AX}\) (\(\text{A} = \text{Sn}, \text{Fe}, \text{Co}, \text{Ni}, \text{Mn}\)). Layered ternary MAX phase materials have the general formula \(\text{M}_{n+1}\text{AX}_n\), where \(\text{M}\) represents a transition metal element, \(\text{A}\) is a group III or IV element of the periodic table, \(\text{X}\) is either \(\text{C}\) or \(\text{N}\), and \(n\) ranges from 1 to 3. It is noted that the MAX phase exhibits the properties of both metals and ceramics. MAX phase compounds such as \(\text{Ti}_3\text{SiC}_2\), \(\text{Ti}_3\text{Au}_2\text{C}_2\), \(\text{Ti}_3\text{SnC}_2\), \(\text{Cr}_2\text{AlC}\), \(\text{Ta}_2\text{AlC}\), and \(\text{Ti}_2\text{Au}_2\text{C}\) were reported to show good tribological properties, high-temperature self-healing ability, and electric conductivity, depending upon the nanolaminated structure of MX slabs with an interlayer of pure \(\text{A}\) element. Examples of the remarkable tribological performance of MAX phase materials were verified by sliding against the tribopair of Ni-based superalloys at an ambient temperature of 550 \(^\circ\)C. The \(\text{Cr}_2\text{AlC}\) MAX phase exhibits excellent wear- and corrosion-resistant properties when used for automotive applications. Further, the incorporation of silver improves the tribological properties significantly and reduces the friction coefficient at room temperature. For example, hot-pressed \(\text{NiMoAl-Cr}_2\text{AlC-Ag}\) composites exhibited a low friction coefficient of 0.27 and a small wear rate, which is attributed to the tribochemical-reacted films of mixed oxides such as \(\text{Ag}_2\text{MoO}_4\) and \(\text{Al}_2\text{O}_3\) formed on the worn surface [88–91]. \(\text{Ti}_3\text{SiC}_2\)-\(\text{PbO-Ag}\) composites fabricated by spark plasma sintering exhibited a friction coefficient of 0.3 and a small wear rate in sliding against Inconel 718 alloy at 800 \(^\circ\)C [92]. Ternary MAX phase materials have also attracted considerable interest as pantograph contact strips for electric railways.

4.4. Chemically Stable Fluorides

Chemically stable fluorides, such as \(\text{CaF}_2\), \(\text{BaF}_2\), and their binary eutectic combinations, are lubricious to reduce friction and wear from about 400 to 900 \(^\circ\)C; however, they exhibit poor tribological behavior at low to moderate temperatures [1,11]. \(\text{CaF}_2\), \(\text{BaF}_2\), and \(\text{BaF}_2-\text{CaF}_2\) eutectics have melting points of 1418 \(^\circ\)C, 1353 \(^\circ\)C, and 1022 \(^\circ\)C, respectively. The lubrication mechanism of alkaline earth fluorides and eutectics depends upon a marked softening effect occurring at temperatures close to 500 \(^\circ\)C due to a brittle-to-ductile transition [1]. Alkaline earth fluorides and eutectics are widely used as solid lubricants in metallic and ceramic matrix composites in industrial applications such as advanced engines due to their high chemical and thermal stability.
CaF$_2$ is a non-layered structure, however, it exists a slip plane of Ca atomic compact plane in its phase structure. Under high-temperature conditions, the slip plane is easy to shear off due to the decrease in atomic force. LiF, MgF$_2$, and some rare-earth fluorides, particularly LaF$_3$ and CeF$_3$, show some lubricating capability in the air at high temperatures up to at least 1000 °C [11]. Most fluorides are able to operate under heavy loads or in chemically reactive environments [1,11]. These solid lubricants can be applied in the form of either coatings or bulk composites, such as fused fluoride coatings or as fluoride/metal or fluoride/ceramic self-lubricating composites or composite coatings. Their coefficients of thermal expansion are matched well with many alloys to which they can be bonded, such as nickel-base superalloys. The tribological properties of rare-earth fluorides and BaF$_2$-CaF$_2$ eutectics at elevated temperatures were investigated [1]. Friction coefficients of nickel-base composite coatings containing CeF$_3$ and LaF$_3$ were from 0.3 to 0.4 with increasing test temperature to about 500 °C. The average coefficient of friction was 0.2 from 500 °C to 1000 °C, which verified the good potential of these fluorides as high-temperature lubricants in oxidizing environments [1]. Interestingly, the relatively high friction of fluoride eutectics at low temperatures and sliding velocities stimulated the synergistic lubrication of combining silver with the BaF$_2$-CaF$_2$ eutectic coatings, which finally led to low and stable friction from room temperature to 900 °C. Sliney pioneered the combination of silver and fluoride eutectics in plasma sprayed composite coatings as a way to reduce friction and wear over a wide temperature range over repeated temperature cycling, especially the development of plasma spray coatings incorporated with silver and BaF$_2$-CaF$_2$ eutectics such as PS212 and PS304 for aerospace applications [7,93]. A particularly important example associated with successful applications of these plasma sprayed coatings is a high-temperature cylinder wall coating for a Stirling engine and a backup lubricant coating for gas bearings [94,95]. Rare-earth fluorides of CeF$_3$, LaF$_3$, and NdF$_3$ were also found to be used as anti-wear, friction-reduction, and extreme pressure additives in lubricating greases and oils [96].

A series of self-lubricating composites based on BaF$_2$-CaF$_2$ eutectic and silver solid lubricants were also developed by powder metallurgy methods [1,94,95]. These composites were designed to consist of a wear-resistant, metal-bonded matrix and solid lubricants of silver and BaF$_2$-CaF$_2$ eutectic. Typical examples of powder metallurgy parts include process valve stem bushings, control surface bearings, and turbine engine bushings [94]. The PM212 composite, which was designed to consist of a metal-bonded Cr$_3$C$_2$ matrix and solid lubricants of Ag and BaF$_2$-CaF$_2$ eutectic, exhibited a friction coefficient of 0.29–0.38 from 25 to 850 °C in the air [95]. Similarly, the PM304 composite, which was comprised of a hardener Cr$_2$O$_3$ within a Ni80Cr20 matrix and Ag and BaF$_2$-CaF$_2$ eutectic solid lubricants, was developed with good tribological performance from room temperature to 650 °C. The friction coefficient of PM304 showed an almost the stable value of 0.33 at temperatures of 200–800 °C [97]. Several Ni$_3$Al matrix composites of Ni$_3$Al-Cr-Ag-BaF$_2$/CaF$_2$ and Ni$_3$Al-Mo-Ag-BaF$_2$/CaF$_2$ were developed based on lubrication characteristics of Ag below 400 °C and fluoride eutectics at elevated temperatures and barium salts formed in situ on the worn surface [98–100].

ZrO$_2$(Y$_2$O$_3$) matrix composites incorporated with BaF$_2$-CaF$_2$ eutectic exhibited good friction and wear properties only above 400 °C when tested in dry sliding against alumina ball in a test temperature range from 25 to 800 °C [101]. In order to generate good self-lubricity from room temperature to about 400 °C, soft noble metals of Au or Ag were added into ZrO$_2$(Y$_2$O$_3$)-CaF$_2$ composites, which finally exhibited low friction coefficients and small wear rates in the order of 10$^{-6}$ mm$^3$/(Nm) from room temperature to 800 °C [102,103].
4.5. Binary Metallic Oxides

Commonly used hard oxides such as alumina, zirconia, and mullite exhibit good wear resistance but generally have high friction coefficients in dry air, which generate high tensile stresses and lead to surface cracks and debris particles [101]. Some oxides have good potential for solid lubrication due to their high thermal stability in the air, even at quite high temperatures. However, they have not been extensively investigated for room temperature solid lubrication due to their typical brittle nature. Their inability to deform plastically or shear easily at room temperature hinders them from forming smooth and lubricious transfer films on worn surfaces. In addition, the wear debris from oxide surfaces is generally abrasive. Oxide surfaces are usually inert in dry air and do not form strong adhesive bonds, such as metals in tribological contact.

In a search for high-temperature solid lubrication, many efforts have been made on various soft oxides. For example, lead monoxide (PbO) has low friction coefficients, especially at elevated temperatures due to its viscous flow, with a friction coefficient of 0.07 at 675 °C. Small amounts of SiO₂, B₂O₃, and Fe₂O₃ are added into PbO to form a sintered ceramic coating that is able to protect against oxidation of PbO and improve adhesion to substrates. Lubricious binary (e.g., PbO, Bi₂O₃, B₂O₃, V₂O₅) and ternary oxides (e.g., Ag₃VO₄, Ag₂MoO₄) are thermally stable and effective lubricants at high temperatures. However, they are generally not capable of providing lubrication at room temperature. These changes in friction and wear properties are closely related to the softening effect after the brittle to ductile transition above a critical temperature. Oxides are known to soften above their ductile-to-brittle transition temperature, which is typically 0.4–0.7 \( T_m \), where \( T_m \) is the melting point (in K).

A crystal-chemical model was proposed to establish the relationship between friction coefficient and ionic potentials of lubricious oxides at sliding interfaces [10]. Re₂O₇, B₂O₃, and V₂O₅ have high ionic potentials and relatively low melting points of 301.5 °C, 450 °C, and 680 °C, respectively. In this case, the higher the ionic potential of oxides, the lower the friction coefficient of binary oxides. Herein, cations are screened more effectively by oxygen anions and thus less likely to interact with other cations at the contact surface, which can be used to explain the decrease in the overall shear strength and the observed lubricious behavior of vanadium or molybdenum oxides. However, some oxides, such as PbO and Bi₂O₃, have very low ionic potentials and exhibit exceptionally low friction, which does not coincide with the above-mentioned model [104].

A polarizability approach was proposed to classify oxides into acidic, basic, and very basic oxides according to the average ionic polarizability, binding energy, optical basicity, and interaction parameter [105]. A relatively low interaction parameter in basic or very basic oxides with large polarizability and low binding energy is closely related to a high unshared electron density and the increased ionicity of chemical bonds [104]. In this case, a high lubricity at elevated temperatures is achieved for an oxide with a low interaction parameter, which has generally a high surface free energy and a low bond strength [106]. Friction for doped oxides or mixtures at the sliding contact is normally reduced by decreasing the interaction parameter. The formation of vacancies and the hopping of ions at the surface of oxides determine to a large extent the frictional behavior at elevated temperatures, which may be valuable to explain the complicated friction behavior of a variety of binary/mixed oxides, based on the polarizability [105].

Computational modeling and simulations are also used to investigate the relationship between crystal structure and frictional behavior of oxides to obtain a better fundamental understanding of experimental observations and make predictions. As is well known, the most widely used atomic-scale modeling approaches are based on density-functional theory (DFT) and molecular dynamics (MD) simulations. DFT-calculations are used to investigate the relationship between layer distance and interlayer bond strength in binary Magneli phases (e.g., V₂O₅, TiO₂, MoO₂, WO₃, and ReO₃), which explains their as-observed low shear behavior by tailoring the chemical composition [107]. However, DFT calculations are generally limited to relatively small systems (on the order of a nanometer) and static
properties, which is quite time-consuming and inefficient. Molecular dynamics simulation deals with the dynamic behavior of moving atoms in successive configurations of a system, which helps to understand friction and wear properties on length scales accessible to experimental methods.

Substoichiometric compounds of certain transition metallic oxides (e.g., Me$_2$O$_{2n-1}$, Me$_x$O$_{3n-1}$, or Me$_{3n}O_{3n-2}$) contain planar lattice faults, which may result in crystallographic shear planes with reduced binding strength. Interestingly, Magneli phases such as TiO$_x$, VO$_x$, MoO$_x$, and WO$_x$ deform by plastic flow rather than by brittle fracture at elevated temperatures [108]. Vanadium-based Magneli phases were verified to possess good self-lubricity in the industrial applications of high-temperature hard nitride coatings up to 700 °C [31]. The friction coefficients are reduced distinctly when these hard coatings were tested from 100 °C to 700 °C until the melting of the V$_2$O$_5$ phase at about 690 °C [109,110]. Hard nitrides or carbides containing W or Mo elements exhibit improved high-temperature tribological properties due to the formation of lubricious Magneli phases such as binary WO$_x$ and MoO$_x$ or multi-component oxides in the sliding contact [111,112]. A variety of binary metallic oxide coatings and their combinations provide solid lubrication at elevated temperatures, but the repeatability and sustainability of solid lubrication are still unknown under long-term thermal cycling conditions. Atomic layer deposited zinc titanate (ZnTiO$_3$) coatings after annealing exhibited a friction coefficient of only 0.12 and a small wear rate of $1 \times 10^{-7}$ mm$^3$/(Nm) in unidirectional sliding against stationary Si$_3$N$_4$ ball, which is potential candidates for high-temperature solid lubrication in moving mechanical assemblies up to 550 °C [28].

In addition, binary or multi-component eutectic oxide systems are of considerable interest for lubricity. As the eutectic points in these systems are normally reduced, they are possible to have lower shear strength than individual oxides. The frictional characteristics of CuO-Re$_2$O$_7$, Cs$_2$O-MoO$_3$, CuO-MoO$_3$, Ag$_2$O-MoO$_3$, PbO-B$_2$O$_3$, PbO-MoO$_3$, CoO-MoO$_3$, Cs$_2$O-SiO$_2$, CuO-V$_2$O$_5$, and NiO-MoO$_3$ systems were studied at different test temperatures to evaluate the effectiveness of two oxide systems [10]. Similarly, the ability to form an easy-to-shear compound or a low-melting-point system can be improved by increasing the difference in ionic potential; therefore, they tend to exhibit lower shear strength and hardness at elevated temperatures [10]. The complex superconductor oxide of YBa$_2$Cu$_3$O$_y$ exhibits a friction coefficient of from 0.20 to 0.50 from room temperature to 1000 °C, which may be considered a potential candidate for high-temperature solid lubricant. YBa$_2$Cu$_3$O$_{7-\delta}$/Ag composites were prepared to evaluate the mechanical and tribological properties from cryogenic to high temperature to identify the contribution of the phonon and electron to friction behavior [113].

Strongly adherent and lubricious thin oxide films can be either directly fabricated by coating technologies, such as reactive magnetron sputtering under an oxygen-containing atmosphere, or by in situ formation by tribo-chemical reaction at elevated temperatures during the wear process. Nowadays, hard coating advancements have evolved from mainly high hardness coatings to current coatings with multifaceted functionalities of high hardness, toughness, temperature stability, oxidation resistance, low friction, and wear [114]. Nitride coatings have generally high hardness and wear resistance, and the vanadium incorporation into the nitride coating contributes to the reduction in the friction coefficient, especially at elevated temperatures via self-adaptation of the hard coating during sliding [31]. Vanadium-containing nitride coatings begin to form lubricious V$_2$O$_5$ on the worn surfaces when the test temperature is increased above 400 °C, and correspondingly, the friction coefficient decreases from 0.45 at room temperature to 0.25 at 700 °C [115]. Similarly, reciprocating wear tests of reactive cathodic arc ion-plated (V, Ti)N coatings were carried out using a ball-on-disc configuration from room temperature to 700 °C. The friction coefficient of (V, Ti)N coatings exhibited a distinct decrease at 500 °C, corresponding to the formation of a reacted layer containing TiO$_2$ and V$_2$O$_5$ oxides. With further increasing the wear test temperature to 700 °C, a melting phenomenon of Magneli phase V$_2$O$_5$ oxides was observed on the worn surface during sliding, which led to a further decrease in friction
coefficient [22]. Likewise, hard nitride coatings form lubricious metallic oxides adaptively on the contact surfaces in high-temperature oxidizing environments [114]. From these studies, V$_2$O$_5$ and related Magneli phases are closely related to the reduction in friction from 450 °C to 650 °C, just below the melting point of V$_2$O$_5$, which is well consistent with the adaptive lubrication mechanisms summarized by Voevodin et al. [30,31]. Some alloys of Ni-Cu-Re, Fe-Re, and Cu-Re exhibit friction coefficients from 0.2 to 0.3 due to the in situ formation of lubricious oxides at elevated temperatures [116]. Considerable progress has been made in identifying and exploring adaptive mechanisms to enable broad-temperature solid lubrication of hard coatings.

Although most binary oxides maintain low shear strengths only in a very narrow temperature range, mainly at elevated temperatures, it is definitely believed that binary oxides have the potential to lubricate in selected environments through suitable surface design and composition tailoring and will become the focus of future studies. The material selected from the oxide family can be operated adaptively at elevated temperatures, in moist air, or in a vacuum.

4.6. Ternary Metallic Oxides

4.6.1. Molybdates

Some metallic molybdates, such as PbMoO$_4$ [117], ZnMoO$_4$, CoMoO$_4$, K$_2$MoO$_4$, BaMoO$_4$ [118], CaMoO$_4$, SrMoO$_4$, Ag$_2$MoO$_4$ and Ag$_2$Mo$_2$O$_7$ [119], are found to be lubricious at elevated temperatures. PbMoO$_4$ and CaMoO$_4$ have melting points of 1070 °C and 1065 °C, and relatively low Mohs hardness of 3.0 and 3.5 at room temperature, respectively. BaMoO$_4$ is one of the most important molybdates with a scheelite-type tetragonal structure and has potential applications in solid lubrication, photoluminescence, solid-state lasers, and photocatalysts. Currently, BaMoO$_4$ and SrMoO$_4$ powders have been prepared by microwave-assisted synthesis, hydrothermal route, micro-emulsion route, complex polymerization method, and electrochemical method [120]. SrMoO$_4$ has a tetragonal structure with the lattice parameters of a = b = 0.539 nm and c = 1.202 nm.

Silver molybdates such as Ag$_2$MoO$_4$ and Ag$_2$Mo$_2$O$_7$ have layered microstructures similar to WS$_2$, which may lead to low friction at high temperatures [27,121]. B-Ag$_2$MoO$_4$ has a typical AB$_2$O$_4$ cubic spinel structure with excellent high-temperature stability, and its related lattice parameters are a = b = c = 0.9318 nm. The unit cell of β-Ag$_2$MoO$_4$ is composed of two basic structural units, with [AgO$_6$] octahedral clusters and [MoO$_4$] tetrahedral clusters. Under the action of external forces, the Ag-O bond is more likely to break and cause interlayer sliding for high-temperature lubrication performance [122]. Table 4 represents the fabrication methods and tribological behavior of self-lubricating materials containing molybdates [117–119,122–124].

From wear tests of Mo$_2$N/MoS$_2$/Ag coatings, the corresponding lubrication mechanism is considered to be associated with the formation of molybdates such as Ag$_2$MoO$_4$ and Ag$_2$Mo$_2$O$_7$ layer sliding during relative motion due to weak bonding [119]. Herein, the combination of silver and molybdenum compounds can provide effective lubrication at low temperatures, while the presence of tribochemical reaction on the worn surface generates lubricious silver molybdates at elevated temperatures. The in situ formation of silver molybdates in plasma sprayed coatings containing silver and pure molybdenum was also studied [123]. As compared with unmodified Ni-based alloy, plasma sprayed coatings exhibited a distinct decrease in both friction and wear at 600 °C and even up to 800 °C. A tribochemically reacted layer was formed on worn surfaces at high temperatures, such as Ag$_2$Mo$_3$O$_{13}$ at 600 °C and Ag$_2$Mo$_2$O$_7$ and Ag$_2$MoO$_4$ at 800 °C, respectively. Especially, the eutectic molybdates of Ag$_2$Mo$_2$O$_7$ and Ag$_2$MoO$_4$, which were melted below 500 °C, were reported [125].
Table 4. Fabrication methods and tribological behavior of self-lubricating materials containing molybdates [117–119,122–124].

| Materials                          | Fabrication Method                  | Tested Conditions                        | Results/Observations                                                                 |
|------------------------------------|-------------------------------------|------------------------------------------|-------------------------------------------------------------------------------------|
| PbMoO$_4$ [117]                    | Pulsed laser deposition              | Ball-on-flat (RT); Pin-on-disc (700 °C)  | • µ: 0.6, poor lubricant and failed immediately (RT)                                        |
|                                    |                                     | 440C steel ball; 1 N; 0.6 m/s; RT, 700 °C | • µ: 0.3–0.4, well lubricated and lasted for over 2 h (700 °C)                           |
|                                    |                                     | Ball-on-disk; Si$_3$N$_4$ ball; load 5 N; 0.126 m/s; RT-600 °C | • µ: 0.6, poor lubricant (RT)                                                        |
|                                    |                                     |                                          | • µ: 0.26, 0.31 (400, 600 °C)                                                          |
|                                    |                                     |                                          | • W: $8 \times 10^{-6}$, $2.15 \times 10^{-5}$, 4.25 $\times 10^{-6}$ mm$^3$/(Nm) (RT, 400, 600 °C) | • The addition of BaMoO$_4$ reduces the friction and wear at high temperature. |
| NiCr-BaMoO$_4$ [118]               | Hot pressing                         | ball-on-disk; Si$_3$N$_4$ ball; load 5 N; 0.11 m/s; RT-600 °C | • µ: 0.6, poor lubricant (RT)                                                        |
|                                    |                                     |                                          | • µ: 0.26, 0.31 (400, 600 °C)                                                          |
|                                    |                                     |                                          | • W: $8 \times 10^{-6}$, $2.15 \times 10^{-5}$, 4.25 $\times 10^{-6}$ mm$^3$/(Nm) (RT, 400, 600 °C) | • The addition of BaMoO$_4$ reduces the friction and wear at high temperature. |
| Mo$_2$N-MoS$_2$-Ag [119]           | Magnetron sputtering deposition      | Ball-on-disk; Si$_3$N$_4$ ball; load 1 N; 0.11 m/s; RT-600 °C | • µ: maximum of 0.48 at 400 °C, minimum of 0.22 at 800 °C |
|                                    |                                     |                                          | • W: 3.2 $\times 10^{-4}$ mm$^3$/(Nm) at 400 °C, minimum of 1.1 $\times 10^{-5}$ mm$^3$/(Nm) at 800 °C | At 1000 °C, abrasive particles affect the formation of Ag$_2$MoO$_4$ lubricant film. |
| NiCoCrAlYTa-Ag-Mo [122]            | High-velocity oxy-fuel spraying      | Ball-on-disk; Al$_2$O$_3$ ball; load 5 N; 0.1 m/s; 25–1000 °C | • µ: maximum of 0.4 at 400 °C, minimum of 0.28 at 800 °C |
|                                    |                                     |                                          | • W: 1 $\times 10^{-4}$ mm$^3$/(Nm) at 400 °C.                                            | • Silver molybdate phases formed at high temperature. |
| NiCrAlY-Ag-Mo [123]                | Atmospheric plasma spraying          | Ball-on-disk; Si$_3$N$_4$ ball; load 5 N; 0.3 m/s; 20–800 °C | • µ: 0.33, 0.28, 0.28, 0.35 (20, 400, 600, 800 °C)                                       |
|                                    |                                     |                                          | • W: 1 $\times 10^{-4}$ mm$^3$/(Nm) at 400 °C.                                            | • Silver molybdate phases formed at high temperature. |
| Ni$_3$Al-Ag-BaMoO$_4$ [124]        | Hot pressing                         | Ball-on-disk; Si$_3$N$_4$ ball; 20 N; 0.19 m/s; 20–800 °C | • µ: maximum of 0.4 at 400 °C, minimum of 0.28 at 800 °C |
|                                    |                                     |                                          | • W: 1 $\times 10^{-4}$ mm$^3$/(Nm) at 400 °C.                                            | • Silver molybdate phases formed at high temperature. |

High-velocity oxy-fuel spraying was used to deposit NiCoCrAlYTa/Ag/Mo composite coatings. A continuous lubricious film comprising layer-like Ag$_2$MoO$_4$ was formed at 800 °C on composite coatings with a friction coefficient of 0.22 and wear rate of 1.1 $\times 10^{-5}$ mm$^3$/(Nm) in sliding against sintered Al$_2$O$_3$ ball [122]. The addition of CaF$_2$ to the Fe-Mo alloys leads to the formation of a surface glaze consisting of CaMoO$_4$, CaF$_2$, Fe$_5$O$_9$, and MoO$_3$ after sliding wear tests at 600 °C. Ternary metal oxides such as Ag-V(Mo,Nb)-O formed on the worn surface are efficient high-temperature solid lubricants due to their structural and chemical inertness, as well as their excellent plasticity and low shear properties. Pulsed laser-deposited PbMoO$_4$ films were lubricious with a friction coefficient of 0.35 and good wear resistance at 700 °C; however, they exhibited a high friction coefficient and failed quickly at room temperature [117].
Nickel-chromium matrix composites incorporated with different amounts of BaMoO$_4$ were hot-pressed to evaluate the tribological properties using a ball-on-disc tribometer up to 600 °C. It is noted that the NiCr-20 wt.% BaMoO$_4$ composite possesses the best tribological performance, with a lower friction coefficient and almost an order of magnitude lower wear rate than the unmodified Ni-Cr composite at 600 °C, which is closely related to the lubricity of a smooth and dense oxide layer formed on worn surfaces, with strong Raman peaks corresponding to BaMoO$_4$ [118]. The self-lubricity of Ni$_3$Al matrix composites in a temperature range of room temperature to 800 °C is attributed to the synergistic lubrication of Ag and BaCrO$_4$ and BaMoO$_3$ barium salts in sliding against coupled Si$_3$N$_4$ ball [124]. However, the formation of non-lubricious BaAl$_2$O$_4$ in the composite must be avoided during the fabrication process.

4.6.2. Tungstates

Some metallic tungstates such as Ag$_2$WO$_4$ [27], ZnWO$_4$ [126,127], CoWO$_4$, CaWO$_4$, BaWO$_4$, and SrWO$_4$ [128] are found to be lubricious at elevated temperatures. CoWO$_4$ has a friction coefficient from 0.2 to 0.25 at temperatures from 600 °C to 800 °C. AWO$_4$ (A = Ca, Ba, and Sr) with a tetragonal scheelite structure is an important material for solid lubrication and electro-optical properties. BaWO$_4$ can be prepared by high-temperature flux crystallization, solid-state reaction, and hydrothermal-electrochemical methods. BaWO$_4$ powders with different morphology, such as olive-like, flake-like, and whisker-like structures, have been successfully prepared through a hydro-thermal process in the presence of different surfactants [129]. Dendrite-like, hollow, and various ordered structures of BaWO$_4$ were prepared by the solution methods, applying organic template and soft hydrothermal conditions, use of polymer or micro-emulsions [130,131]. SrWO$_4$ exhibits a tetragonal structure and has lattice parameters of $a = b = 0.542$ nm and $c = 1.195$ nm. In addition, the tribological performance of Ag$_2$WO$_4$ at elevated temperatures was investigated by means of ab initio calculations through MD simulations [27].

Ni$_3$Al-based composites incorporated with silver, BaF$_2$-CaF$_2$, and W were produced using powder metallurgy to evaluate tribological performance up to 800 °C on a ball-on-disc high-temperature tribometer. Stable friction and low wear rates are attributed to the formation of CaWO$_4$ and BaWO$_4$ during sliding at elevated temperatures [128].

Thin WS$_2$-ZnO composite films are deposited to produce lubrication over a wide temperature range. The tungsten disulfide reacted with the zinc oxide to form a lubricious ZnWO$_4$ phase at elevated temperatures [126]. Similarly, tribological properties of burnished films comprising WS$_2$ and ZnO are tested at high temperatures using a unidirectional ball-on-disc friction and wear tester [127]. The lubricious ternary ZnWO$_4$ oxide layer formed at 500 °C leads to low and stable friction during wear tests. These composite coatings were adaptively lubricious where WS$_2$ provided the low-temperature lubrication and then reacted with zinc oxide to form an effective high-temperature lubricant. However, the drawback associated with this synergistic lubrication is that the low-temperature lubrication of WS$_2$ will lose inevitably due to an irreversible reaction once returning to room temperature during the long-term thermal cycling process.

4.6.3. Vanadates

Lubricious binary V$_2$O$_5$ and ternary oxides of AgVO$_3$ [132,133], Ag$_3$VO$_4$ [27,121], BiVO$_4$, Bi$_4$V$_2$O$_{11}$, and Bi$_2$V$_3$O$_9$ are thermally stable and effective lubricants at high temperatures, although they are generally ineffective at providing self-lubricity at room temperature. Vanadate is a pentavalent monomer of vanadium oxide that can exist either as the meta- or ortho- form depending on the number of oxygen ligands around the vanadium atom. Vanadate powders with different particle size distributions and morphologies have been synthesized by simple and facile approaches such as hydrothermal and wet precipitation routes at room temperature. BiVO$_4$ powders with a monoclinic structure were also synthesized by a high-temperature solid-state reaction at 700 °C. Many efforts have been made on the synthesis and characterization of monoclinic phase β-AgVO$_3$ and
nanostructured Ag@Ag₃VO₄ [134,135]. Silver vanadates such as Ag₃VO₄ have layered microstructures, which may lead to low friction at high temperatures [27,121]. Table 5 shows the fabrication methods and tribological behavior of self-lubricating materials containing vanadates [121,132,133].

Table 5. Fabrication methods and tribological behavior of self-lubricating materials containing vanadates [121,132,133].

| Materials                  | Fabrication Method                  | Tested Conditions | Results/Observation                                                                 |
|----------------------------|-------------------------------------|-------------------|-------------------------------------------------------------------------------------|
| VN-Ag [121]                | Magnetron sputtering deposition     | Ball-on-disk; Si₃N₄ ball; load 2 N; 0.11 m/s; RT-1000 °C | • μ: 0.37, 0.3, 0.12, 0.2 (RT, 350, 700, 1000 °C)                                           |
|                            |                                     |                   | • Silver, vanadium oxide and silver vanadate formed on the surface.                  |
|                            |                                     |                   | • Ag₃VO₄ with layered atomic structure is presumed to improve the lubricant performance. |
| VN-Ag [132]                | Pulsed laser deposition             | Ball-on-disk; Al₂O₃ ball; load 10 N; 0.063 m/s; RT-900 °C | • μ: 0.3, 0.2, 0.18, 0.1, 0.08 (RT, 200, 400, 600, 800 °C)                                |
|                            |                                     |                   | • The synergy of silver vanadate and molten lubricant vanadium oxides generates a low friction coefficient at high temperature. |
| NiCrAlY-Cr₃C₂(NiCr)-V₂O₅-Ag₂O [133] | Laser cladding                  | Ball-on-disk; Si₃N₄ ball; load 3 N; 0.188 m/s; 25–800 °C | • μ: 0.52, 0.49, 0.35, 0.25, 0.15 (25, 350, 600 °C)                                           |
|                            |                                     |                   | • W: maximum of 6.9 × 10⁻⁵ mm³/(Nm) at 400 °C, minimum of 2.9 × 10⁻⁵ mm³/(Nm) at 800 °C |

Silver-based vanadate (Ag₀.₃₃V₂O₅) thin films deposited by PVD have a higher friction coefficient at moderate temperatures but exhibit a lower friction coefficient up to 500 °C and even to about 0.25 at 600 °C than V₂O₅ coatings [109]. VN thin films incorporated with different silver contents deposited by unbalanced reactive magnetron sputtering exhibit good frictional behavior within the temperature ranges up to 1000 °C using a ball-on-disc friction and wear tester, with a minimum friction coefficient of 0.2 at 700 °C, which is attributed to the in situ formation of lubricious silver vanadate an Ag₃VO₄ during sliding [121]. Similarly, Ag/VN thin films by pulsed-laser deposition also exhibited reduced friction and wear at 700 °C and 900 °C when sliding against an Al₂O₃ ball, due to the formation of V₂O₅, AgVO₃, and Ag₃VO₄ [132]. Laser-clad NiCrAlY-based coatings incorporated with V₂O₅ and Ag₂O solid lubricants have low friction and wear from 600 °C to 800 °C, due to the formation of AgVO₃ and Ag₃VO₄ [133]. In addition, soft metal-containing inorganic compounds such as Ag₃VO₄ were also embedded into textured hard surfaces to produce continuous self-lubricity over a wide temperature range up to 800 °C [136].

4.6.4. Tantalates

Ternary metal oxides of (Ag, Cu)-(Ta(V,Mo))-O such as AgTaO₃, CuTaO₃, and CuTa₂O₆ have recently been explored as promising lubricants for high-temperature applications due to their structural and chemical inertness and low friction at elevated temperatures. A soft metallic silver-containing phase is generated when silver tantalate (AgTaO₃) with a layer-like structure is highly resistant to oxidation and is subjected to sliding at elevated temperatures [137]. Layered AgTaO₃ has a relatively high melting temperature of 1172 °C and undergoes a series of structural phase transitions with temperature until it reaches its melting point [138]. The temperature dependence of mechanical and tribological responses to sliding is closely linked with the above-mentioned structural changes of AgTaO₃.
Several processing approaches were explored to fabricate lubricious silver tantalate films on Inconel substrates for extreme temperature applications as follows: (1) burnished powder films on the substrate; (2) magnetron sputtered monolithic silver tantalate films; (3) sputter-deposited adaptive tantalum nitride/silver nanocomposite coatings, which forms a lubricious silver tantalate layer on its surface by high-temperature tribochemical reaction during sliding. For example, this coating has friction coefficients in the range from 0.06 to 0.15 at 750 °C in dry sliding against Si3N4 counterfaces [137,139]. A mechanically mixed layer comprising nanocrystalline Ag, Ta2O5, and AgTaO3 phases was generated due to surface reconstruction induced by friction heat and shear stress. The aggregated silver clusters on the sliding surface are beneficial for low friction [139]. However, the silver diffuses rapidly on the surface, and its high mobility leads to a relatively low wear resistance for AgTaO3. It is more noteworthy that silver particle clusters were observed to migrate randomly and finally led to system failure in various sliding components at elevated temperatures [139–141].

Recently, density-functional theory (DFT) and molecular dynamics (MD) simulations have been used to reveal the friction and wear processes of three ternary oxides—AgTaO3, CuTaO3, and CuTa2O6—through newly developed empirical potential parameters and experimental measurements. The sliding mechanisms are verified through both experimental characterization of the film composition before and after sliding, and quantification of Ag or Cu cluster formation during the evolution of the film in MD, and DFT energy barriers for atomic migration on the surface. All the experimental observations and theoretical calculations supported the hypothesis that metal (or metal oxide) clusters generated on the sliding surface exert a strong influence on friction and wear behavior [140]. The friction and wear performance of AgTaO3 at elevated temperatures were investigated through ab initio MD simulations [141], which also quantified the presence and migration of silver clusters by contact sliding simulations [139].

4.6.5. Alkaline Earth Metallic Chromates

In recent years, ternary chromate compounds of MCrO4, MCr2O4, and MCrO3 (M = Ba, Sr, and Ca) between chromium sesquioxide and alkaline earth metallic oxides have been investigated as potential solid lubricants for self-lubricating metallic or ceramic matrix composites [15,142–144]. In the MCrO4 class of oxometallates, M represents different alkaline earth metals with oxidation states of +2. BaCrO4 consists of [CrO4]2− tetrahedra and the coordinated Ba2+ cations, and each Ba2+ cation connects with 7 tetrahedra of [CrO4]2− with a coordination number of 12. Figure 3 shows the schematic diagram of the crystal structure of BaCrO4. BaCrO4 has an orthorhombic structure with lattice parameters of a = 0.9113 nm, b = 0.5528 nm, and c = 0.7336 nm. The first principle calculations and the Castep modules are used to calculate the bulk modulus and compression coefficient of BaCrO4, which are 28.1GPa and 0.0357 GPa−1, respectively. BaCrO4 has been widely used as a model system for the study of morphological control of inorganic minerals, as well as an oxidizing agent, a catalyst for enhancing vapor-phase oxidation reactions, and high-temperature solid lubricants [15,145].

Oxides of some bivalent metallic elements and trivalent chromium form a group of oxometallates, which have the general formula of M2+Cr3+O42− and exhibit two different crystal structures depending upon the ionic radius ratio of M2+/Cr3+. Most MCr2O4-type (M = Fe, Mn, Cu, Mg, Zn, Ni, and Co) compounds with a spinel structure have good thermal stability. However, alkaline earth metallic oxometallates such as BaCr2O4, SrCr2O4, and CaCr2O4 are another part of the family of M2+Cr3+O42− compounds [146]. These chromates have a layered structure with triangular sheets of formula CrO2 separated by M2+ in trigonal prismatic coordination [142]. Figure 4 shows the schematic diagram of the crystal structure of MCr2O4 (M = Ca, Ba, Sr) [142]. BaCr2O4 is an orthorhombic phase with lattice parameters of a = 1.2286 nm, b = 0.5921 nm, and c = 0.5146 nm, and is thermally stable in the non-oxidizing atmosphere of argon. It does not decompose up to 1400 °C.
The calculated bulk modulus and compression coefficient of BaCr$_2$O$_4$ are 71.6 GPa and 0.014 GPa$^{-1}$, respectively.

![Schematic diagram of crystal structure of BaCrO$_4$.](image1)

**Figure 3.** Schematic diagram of crystal structure of BaCrO$_4$.

![Schematic diagram of crystal structure of MCr$_2$O$_4$ (M = Ca, Ba, Sr).](image2)

**Figure 4.** Schematic diagram of crystal structure of MCr$_2$O$_4$ (M = Ca, Ba, Sr). Reproduced with permission from Reference [142], Copyright © Springer 2012.

BaCrO$_3$ exhibits a variety of layered structures with 4, 6, 12, 14, and 27 layers, which correspond to various c/a values of 1.654, 2.433, 4.901, 5.752, and 11.101, respectively. The local density approximation (LDA) method is used to calculate the properties of three kinds of hexagonal BaCrO$_3$ structures (e.g., P63/mmc (194, Z = 4, 6, and 14)), which have bulk moduli of 94.7 GPa, 100 GPa, and 85.5 GPa, and compression coefficients of 0.0106 GPa$^{-1}$, 0.01 GPa$^{-1}$ and 0.0117 GPa$^{-1}$, respectively.

BaCrO$_4$ particles with various size distributions and morphologies have been synthesized in different synthesis routes in the open literature. For example, BaCrO$_4$ powders with different crystallographic morphologies and sizes were prepared by the chemical co-precipitation method, which is important for large-scale production when considered as a high-temperature solid lubricant [145]. The influences of temperature and atmosphere on the thermal stability of BaCrO$_4$ powders synthesized by a facile aqueous solution route were evaluated. Differential thermal analysis-thermogravimetry (DTA-TG) and X-ray
wear process [142]. In a previous study, BaCrO₄ is not thermally stable during vacuum sintering because the sample containing BaCrO₄ changes its color from yellow to green on polished surfaces after vacuum sintering at 1050 °C [143]. Generally, thermal stability must be taken into account if the lubricant is designed to operate at elevated temperatures and different atmospheres. Gontarz [148] claimed that BaCrO₄ is decomposed to a mixture of Ba₂CrO₄ and BaCr₂O₄ with the release of oxygen. Similarly, Azad et al. [149] also reported that BaCrO₄ is decomposed into BaCr₂O₄ and BaO-Cr₂O₃-CrO₃ beyond 900 °C in the non-oxidizing atmosphere according to the Ba-Cr-O phase diagram.

Microsized BaCr₂O₄ powders were synthesized by a solid-state reaction method using stoichiometric amounts of BaCO₃ and Cr₂O₃ powders calcined at 1250 °C for 3 h in a vacuum [150]. In a previous study, BaCr₂O₄ was found to be thermally unstable in an air atmosphere at elevated temperatures. An oxidation reaction of BaCr₂O₄ occurs at 790.2 °C, which leads to the oxidized products of BaCrO₄ and Cr₂O₃. It is supposed that such an oxidation reaction contributes to the self-lubricity at elevated temperatures during wear tests. The formed BaCrO₄ lubrication layer is easy-to-shear and is prone to spread out in the wear track at elevated temperatures [142].

A ball-on-disc high-temperature friction and wear tester was used to investigate the friction and wear characteristics of pure BaCr₂O₄ ceramics. They were investigated up to 800 °C in dry sliding against an alumina ball. BaCr₂O₄ crystallizes with connections between the layers of edge-shared CrO₆-octahedra and the [BaO₄]-chains, which is also beneficial for reducing friction and wear. BaCr₂O₄ ceramics have low friction coefficients and small wear rates at temperatures increasing up to 400–600 °C. Interestingly, BaCr₂O₄ is oxidized in air to form a smooth self-lubricating film comprising BaCrO₄ and Cr₂O₃ on the worn surface to effectively reduce friction and wear. However, if severe oxidation leads to a decrease in the relative density of pure BaCr₂O₄ ceramics, it further expedites the wear process [142].

Table 6 shows the fabrication methods and tribological behavior of self-lubricating materials containing alkaline earth metallic chromates [15,142–144,150,151]. Both BaCrO₄ and BaCr₂O₄ can be added into a metallic or ceramic matrix to fabricate self-lubricating composites or coatings by a variety of fabrication technologies such as powder metallurgy [143,151], low-pressure plasma spraying [15], and electrodeposition methods [150]. For example, the spark-plasma-sintered ZrO₂(Y₂O₃)-50BaCrO₄ composite after 800 °C wear test in sliding against alumina ball is shown in Figure 5. Clearly, the barium chromate becomes softening at 800 °C, which spreads out on the sliding surfaces under tribo-stressing and creates a self-lubricating fine grain layer. The lubrication mechanism is associated with the in situ formation of surface glaze with ultrafine nanograins due to thermo-mechanical induced recrystallization/deformation at elevated temperatures. In this case, grain boundary sliding and grain rotation of BaCrO₄ nanograins in the glaze layer also promote plastic smearing and self-lubricity at elevated temperatures. Low-pressure plasma sprayed ZrO₂-BaCrO₄ coating also showed good lubricity at elevated temperatures [15].
Table 6. Fabrication methods and tribological behavior of self-lubricating materials containing alkaline earth metallic chromates [15,142–144,150,151].

| Materials                  | Fabrication Method          | Tested Conditions                                                                 | Results/Observations                                                                                                                                 |
|----------------------------|----------------------------|-----------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|
| ZrO$_2$(Y$_2$O$_3$)-BaCrO$_4$ [15] | Low-pressure plasma spraying | Ball-on-block; Al$_2$O$_3$ ball; load 50 N; frequency of 10 Hz with stroke of 1 mm; RT-800 °C | • $\mu$: 0.81, 1, 0.92, 0.58, 0.57, 0.5 (RT, 100, 200, 400, 600, 800 °C);  
  • Wear depth: 42% reduction from 100 °C to 800 °C;  
  • Lubrication resulting from the formation of transfer BaCrO$_4$ films at high temperatures.  
  • $\mu$: 0.7, 0.38, 0.35, 0.4 (RT, 400, 600, 800 °C)  
  • $W$: $9.7 \times 10^{-5}$, $1.56 \times 10^{-5}$, $1.43 \times 10^{-5}$, $4.9 \times 10^{-5}$ mm$^3$/(Nm) (RT, 400, 600, 800 °C)  
  • Friction heat at the surface contributes to the oxidization of BaCr$_2$O$_4$ at between 400 and 600 °C.  
  • BaCrO$_4$ as oxidized product dominates the formation of lubricating films.  
  • Severe oxidation at 800 °C degrades the wear resistance.  
  • Addition of 50 wt.% BaCrO$_4$ significantly improves tribological properties at high temperatures.  
  • $\mu$: 0.65, 0.81, 0.6, 0.7, 0.55 (RT, 200, 400, 600, 800 °C)  
  • $W$: $6.5 \times 10^{-5}$, $3 \times 10^{-5}$, $1.6 \times 10^{-5}$, $3 \times 10^{-5}$ mm$^3$/(Nm) (RT, 200, 400, 600, 800 °C)  
  • Sintered samples contain a lot of pores.  
  • $\mu$: 0.6, 0.7, 0.8, 0.7, 0.6 (RT, 200, 400, 600, 800 °C)  
  • $W$: $3.1 \times 10^{-5}$, $2.2 \times 10^{-5}$, $3 \times 10^{-6}$, $4 \times 10^{-6}$ mm$^3$/(Nm) (RT, 200, 400, 600, 800 °C)  
  • SiO$_2$ doping as sintering additive improves the density of composites, resulting in notably lower wear rate, especially at high temperatures.  
  • $\mu$: 0.7, 1.2, 0.44, 0.46, 0.5 (RT, 200, 400, 600, 800 °C)  
  • $W$: $5 \times 10^{-5}$, $6 \times 10^{-5}$, $2 \times 10^{-6}$, $2 \times 10^{-6}$, $1.2 \times 10^{-5}$ mm$^3$/(Nm) (RT, 200, 400, 600, 800 °C)  
  • Silver instead of BaCrO$_4$ works as lubricant at 600 °C. |
Table 6. Cont.

| Materials Fabrication Method Tested Conditions Results/Observations |
|---------------------------------------------------------------|
| NiCr-BaCr₂O₄ [151] Hot pressing Ball-on-disk | \( \mu \) 0.72, 0.27, 0.25, 0.27 (RT, 400, 600, 800 °C)  \( W \) \( 2.29 \times 10^{-5}, 9 \times 10^{-6}, 2.6 \times 10^{-6} \) \( 4.5 \times 10^{-6} \text{ mm}^3/(\text{Nm}) \) (RT, 400, 600, 800 °C)  Addition of BaCr₂O₄ significantly reduces friction and wear of NiCr alloy at 400–800 °C.  \( \mu \) 0.31  \( W \) 2.79 \( \times 10^{-6} \text{ mm}^3/(\text{Nm}) \) (RT)  Enhanced microhardness 745HV  BaCr₂O₄ particles enhance the hardness and the resistance to scuffing wear of composite coating. |
| Ni-16.6 vol.% BaCr₂O₄ [150] Electrodeposition Ball-on-disk | \( \mu \) 0.31  \( W \) 2.79 \( \times 10^{-6} \text{ mm}^3/(\text{Nm}) \) (RT)  Enhanced microhardness 745HV  BaCr₂O₄ particles enhance the hardness and the resistance to scuffing wear of composite coating. |

Figure 5. Worn surface of ZrO₂(Y₂O₃)-50BaCrO₄ composite after 800 °C wear test: (a) morphology, (b) enlarged view of (a). Reproduced with permission from Reference [143], Copyright © Elsevier B.V. 2005.

4.7. Alkaline Earth Metallic Sulfates

Alkaline earth metallic sulfates, such as BaSO₄ (barite), SrSO₄ (celestite), and CaSO₄ (anhydrite), have been widely used as important inorganic chemical products in ceramics, pigments, electronics, cosmetics, papermaking, and high-temperature solid lubricants [152,153]. Especially the excellent lubricating properties of barite and celestite as well as their sulfate solid solutions are closely related to their structures and morphologies. The lubrication mechanisms are associated with the slippage along the basic plane of (001) at room temperature and the in situ formation of surface glaze with ultrafine nanograins due to thermo-mechanical induced recrystallization/deformation at elevated temperatures. Similar to the BaCrO₄ crystal, SrSO₄ consists of \([\text{SO}_4]^{2-}\) tetrahedra and the coordinated Sr\(^{2+}\) cations, and each Sr\(^{2+}\) cation connects with 7 tetrahedra of \([\text{SO}_4]^{2-}\) with a coordination number of 12. Figure 6 shows the schematic diagram of atomic arrangements at planes of (002) and (210) in the SrSO₄ crystal. Alkaline earth sulfate particles with a variety of unusual and well-defined crystallographic morphologies were obtained via controlled nucleation and growth by using various additives as crystal modifiers from an aqueous solution [154]. SrSO₄ has an orthorhombic structure with lattice parameters of \( a = 0.8359 \) nm, \( b = 0.5352 \) nm, and \( c = 0.6866 \) nm, while BaSO₄ has the same structure with lattice parameters of \( a = 0.8881 \) nm, \( b = 0.5454 \) nm, and \( c = 0.7157 \) nm at room temperature. TG/DTA thermal analysis indicates that barite and celestite as well as their sulfate solid solutions are thermally stable, and have almost no weight loss up to 1300 °C and only a structural transition of orthorhombic to monoclinic phase at about 1100 °C. The formation process of
sulfate hierarchical architectures is quite complicated and is affected by crystal structure and crystal growth environments, such as surface energy, the degree of supersaturation, and diffusion of the reaction [152–155].

A facile aqueous solution route was employed to prepare SrSO$_4$ nanocrystals from a needle-like to a tablet-like feature by increasing the molar ratio (R) of Sr$^{2+}$ cations to SO$_4^{2-}$ anions from R = 1:2 to R = 2:1 at room temperature without using any surfactants or templates [152]. The crystallographic morphology of SrSO$_4$ depends mainly upon the preferential absorption of excessive strontium cations on the (020) and (210) planes. Monodispersed peanut-type SrSO$_4$ particles with an average length of 1.7 µm and an aspect ratio of 1.4 were synthesized using a facile precipitation reaction of Sr-EDTA chelating precursors at ambient temperature. These peanut-type SrSO$_4$ particles have a relatively large Brunauer–Emmett–Teller (BET) surface area of about 20.9 m$^2$ g$^{-1}$ and contain mesopores with a mean pore size of about 34.3 nm [153]. The Ba$_x$Sr$_{1-x}$SO$_4$ solid solution nanocrystals synthesized by the chemical precipitation method have an orthorhombic structure, an average size of 80–100 nm, and an ellipsoidal-shaped morphology. These Ba$_x$Sr$_{1-x}$SO$_4$ solid solutions with different composition parameters are perfectly indexed as a single phase of orthorhombic with a space group, Pbnm (62) of Ba$_{0.25}$Sr$_{0.75}$SO$_4$, Ba$_{0.5}$Sr$_{0.5}$SO$_4$, and Ba$_{0.75}$Sr$_{0.25}$SO$_4$, respectively [155].

Fabrication methods and tribological behavior of self-lubricating materials containing alkaline earth sulfates are summarized in Table 7 [101,136,156–160]. In our previous work, the alkaline earth sulfates with a barite-type structure such as BaSO$_4$, SrSO$_4$, and Ba$_x$Sr$_{1-x}$SO$_4$ possessed excellent self-lubricity, thermo-chemistry stability, and innocuity. These sulfates are also verified to possess lubrication ability, especially at elevated temperatures. Textured hard surfaces embedded with BaSO$_4$ additives are able to provide sustainable self-lubricity [136]. In fact, barite exhibits good tribological properties, which only change slightly with the increase in temperature, and the wide availability of the barite mineral greatly reduces the cost. Barite can be used as the lubricant filler for brake pads. Both the coefficient of friction and wear rates are reduced by increasing the barite ratio in a brake pad. Interestingly, the variations in sliding velocity and rubbing pressure have little effect on the friction coefficient of the barite-containing brake pad materials. High stability of the friction coefficient and excellent fade resistance are guaranteed by increasing the BaSO$_4$ content in the friction material. Particularly, barite is resistant to high temperatures and has no significant change in its properties at temperatures above 300 °C [161,162].
Table 7. Fabrication methods and tribological behavior of self-lubricating materials containing alkaline earth sulfates [101,136,156–160].

| Materials | Fabrication Method | Tested Conditions | Results/Observations |
|-----------|-------------------|------------------|---------------------|
| \(\text{Al}_2\text{O}_3\)-Mo-BaSO\(_4\) [136] | BaSO\(_4\) was burnished onto textured surface of hot pressed \(\text{Al}_2\text{O}_3\)-Mo. | Pin-on-disk; \(\text{Al}_2\text{O}_3\) pin; load 70 N; frequency of 10 Hz with stroke of 1 mm; RT-800 °C | ● \(\mu\): 0.45, 0.57, 0.44, 0.3, 0.3 (RT, 200, 400, 600, 800 °C)  
● BaSO\(_4\) effectively improves the wear resistance and could remain a lot in micro-dimples. |
| \(\text{ZrO}_2\)(Y\(_2\)O\(_3\))-BaSO\(_4\) [101] | Spark plasma sintering | Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 5 N; frequency of 1 Hz with stroke of 10 mm; RT-800 °C | ● \(\mu\): 0.36, 0.3, 0.33 (RT, 400, 800 °C)  
● \(W\): \(4.72 \times 10^{-6}\) mm\(^3\)/(Nm) (800 °C)  
● Softened BaSO\(_4\) fine-grain layer forms on the worn surfaces at high temperature. |
| \(\text{Al}_2\text{O}_3\)-SrSO\(_4\) [156] | Spark plasma sintering | Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 3 N; frequency of 1 Hz with stroke of 10 mm; RT-800 °C | ● \(\mu\): 0.38, 0.44, 0.42, 0.35, 0.29 (RT, 200, 400, 600, 800 °C)  
● \(W\): maximum of \(8 \times 10^{-4}\) mm\(^3\)/(Nm) at 200 °C, minimum of \(1 \times 10^{-5}\) mm\(^3\)/(Nm) at 600 °C |
| \(\text{Al}_2\text{O}_3\)-PbSO\(_4\)-SiO\(_2\) [156] | Spark plasma sintering | Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 5 N; frequency of 1 Hz with stroke of 10 mm; RT-800 °C | ● \(\mu\): 0.32, 0.36, 0.26, 0.26, 0.31 (RT, 200, 400, 600, 800 °C)  
● \(W\): maximum of \(1 \times 10^{-4}\) mm\(^3\)/(Nm) at 800 °C, minimum of \(1 \times 10^{-5}\) mm\(^3\)/(Nm) at 200 °C |
| \(\text{Al}_2\text{O}_3\)-BaSO\(_4\)-Ag [156] | Spark plasma sintering | Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 3 N; frequency of 1 Hz with stroke of 10 mm; RT-800 °C | ● \(\mu\): 0.38, 0.3, 0.21, 0.21, 0.22 (RT, 200, 400, 600, 800 °C)  
● \(W\): maximum of \(4 \times 10^{-4}\) mm\(^3\)/(Nm) at 800 °C, minimum of \(5 \times 10^{-6}\) mm\(^3\)/(Nm) at 200 °C |
| \(\text{ZrO}_2\)(Y\(_2\)O\(_3\))-Al\(_2\)O\(_3\)-Ba\(_{0.5}\)Sr\(_{0.5}\)SO\(_4\) [157] | Spark plasma sintering | Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 5 N; frequency of 1 Hz with stroke of 10 mm; RT, 760 °C | ● \(\mu\): 0.21, 0.11 (RT, 760 °C)  
● \(W\): \(5 \times 10^{-5}\), \(3.6 \times 10^{-6}\) mm\(^3\)/(Nm) (RT, 760 °C)  
● Friction coefficient at 760 °C is much more stable than that of unmodified \(\text{ZrO}_2\)(Y\(_2\)O\(_3\))-Al\(_2\)O\(_3\) composites due to the formation and effective spreading of Ba\(_{0.5}\)Sr\(_{0.5}\)SO\(_4\) lubricating film. |
| Fe\(_3\)Al-Ba\(_{0.25}\)Sr\(_{0.75}\)SO\(_4\) [158] | Hot pressing | Ball-on-disk; Si\(_3\)N\(_4\) ball; load 10 N; \(0.01\) mm/s; RT-800 °C | ● \(\mu\): 0.3, 0.6, 0.54, 0.19, 0.29 (RT, 200, 400, 600, 800 °C)  
● \(W\): maximum of \(4 \times 10^{-5}\) mm\(^3\)/(Nm) at 400 °C  
● The main wear mechanism is abrasion wear at high temperature. |
| Ni-6.83 vol.% SrSO\(_4\) [159] | Electrodeposition | Ball-on-disk; SAE52100 bearing ball; load 0.5 N; rotating speed 50 rpm; rotating radius 5mm; RT Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 5 N; frequency of 1 Hz with stroke of 10 mm; RT-800 °C | ● \(\mu\): 0.18, 0.2, 0.22, 0.26, 0.27 (RT, 200, 400, 600, 800 °C)  
● \(W\): \(< 2 \times 10^{-6}\) mm\(^3\)/(Nm) |
| SrSO\(_4\)-Ag [160] | Chemical precipitation | Ball-on-disk; SAE52100 bearing ball; load 0.5 N; rotating speed 50 rpm; rotating radius 5mm; RT Ball-on-block; \(\text{Al}_2\text{O}_3\) ball; load 5 N; frequency of 1 Hz with stroke of 10 mm; RT-800 °C | ● \(\mu\): 0.18, 0.2, 0.22, 0.26, 0.27 (RT, 200, 400, 600, 800 °C)  
● \(W\): \(< 2 \times 10^{-6}\) mm\(^3\)/(Nm) |
Various approaches were developed to fabricate the self-lubricating sulfate-containing composites, such as plasma spraying, physical vapor deposition, electrodeposition, hot pressing, spark plasma sintering, and so forth. BaSO₄, SrSO₄, and (Ba,Sr)SO₄ can be added or in situ formed into a metallic or ceramic matrix to synthesize self-lubricating composites or coatings by powder metallurgy [101,156–158,163], Pulsed laser deposition [126], electrodeposition [159], and burnishing methods [160,164]. For example, the friction and wear properties of ZrO₂(Y₂O₃)-Al₂O₃-50BaSO₄ composites are distinctly improved at elevated temperatures, as contrasted with unmodified ZrO₂(Y₂O₃)-Al₂O₃ ceramics. Figure 7 shows the frictional behavior of the ZrO₂(Y₂O₃)-Al₂O₃-50BaSO₄ composite as a function of the wear cycle at different temperatures when sliding against an alumina ball. BaSO₄-containing composite has a friction coefficient of 0.33 and a wear rate of 4.72 × 10⁻⁶ mm³/(Nm) at 800 °C. Figure 8 shows the worn surface of the ZrO₂(Y₂O₃)-Al₂O₃-50BaSO₄ composite after the 800 °C wear test. Clearly, a self-lubricating BaSO₄ fine-grain layer is observed on the sliding surfaces at elevated temperatures, which avoids the direct tribo-contact between the oxide ceramics and the coupled ball. The lubrication mechanisms at elevated temperatures are associated with the formation of surface glaze with ultrafine nanograins due to thermo-mechanical induced recrystallization/deformation. Similarly, grain boundary sliding and grain rotation of BaSO₄ nanograins in the glaze layer also promote plastic smearing and self-lubricity at elevated temperatures [101].

Figure 7. Friction coefficients of ZrO₂(Y₂O₃)-Al₂O₃-50BaSO₄ composite as a function of wear cycle at different temperatures.

Figure 8. Worn surface of ZrO₂(Y₂O₃)-Al₂O₃-50BaSO₄ composite after 800 °C wear test: (a) morphology, (b) enlarged view of (a). Reproduced with permission from Reference [101], Copyright © Elsevier B.V. 2009.
Alkaline earth sulfate-containing coatings also contribute to good tribological properties. Low friction coefficients and small wear rates are validated for SrSO$_4$ or SrSO$_4$-Ag films coated on ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$ and Si$_3$N$_4$ ceramics by the chemical precipitation method over a wide temperature range [160,164]. Pulsed laser-deposited CaSO$_4$-containing films tend to be fairly soft and to be easily deformed, which lubricates better at elevated temperatures than CaF$_2$ when incorporated into a layered structure with silver [126]. The friction and wear properties of BaSO$_4$ and SrSO$_4$ powder films coated on different substrates of Al$_2$O$_3$ and SUS316 stainless steel are investigated by using a reciprocating ball-on-block high-frequency induction-heated tribometer to slide against Al$_2$O$_3$ or SUS316 steel balls, respectively. Both the BaSO$_4$ and SrSO$_4$ powder films coated on Al$_2$O$_3$ substrate by chemical precipitation have average friction coefficients of less than 0.4 up to 800 °C in air. In particular, the friction coefficients of flake-shaped BaSO$_4$ powder films coated on an Al$_2$O$_3$ substrate are lower than those of lump-shaped BaSO$_4$ powder films. The BaSO$_4$-10mass% Ag films formed on the SUS316 substrate have average friction coefficients from 0.2 to 0.4 from room temperature to 800 °C in the air [164].

4.8. Silicates

The silicates generally have the building blocks of (SiO$_4$)$_{4-}$ units, where each O$^{2-}$ ion is coordinated with two Si$^{4+}$ ions, so the tetrahedral join at corners. The SiO$_4$ tetrahedra can be either kept separated or linked to one another to form chains or rings, which can be made into sheets or double chains. A special feature of silicates is that the cations that are outside the SiO$_4$ tetrahedra are quite easy to be replaced, e.g., the replacement of Si$^{4+}$ with other similar-sized ions such as Al$^{3+}$ without changing the nature of oxygen coordination. Layered micas are very rigid but cleave very easily along a plane. The bonds within the layers are very strong, but the van der Waals bonds between the layers are weak. Therefore, micas and other minerals such as serpentine and attapulgite are designed to supply easy paths for crack propagation and improve the machinability of some commercial ceramics [165].

Inorganic silicate polymers are salts of the oxyacids of silicon. The silicate anion structure is formed by repeated condensation of tetrahedral SiO$_4^{4-}$ groups, by linking through the covalent bridging oxygen atoms to form chains, cyclic and larger polymeric structures [166]. Silicate-based additives have been used in the lubrication of bearings or gearboxes under heavy loads or high-temperature circumstances. The sodium or potassium silicate reacts with metallic rubbing surfaces to form a lubricating film of metallic silicate on the sliding interface. Silicate-based compounds such as Al$_4$(Si$_4$O$_{10}$)(OH)$_4$ [167], and aluminum–magnesium silicate [168] are designed to protect the surfaces of engines and reduce effectively friction and wear as lubrious additives. The silicates have self-restoration capability at the sliding contacts [169]. These additives can self-repair the mechanical damage at the sliding surfaces, as evidenced by the mineral elements within the tribofilm, which are different from the base materials. However, the silicates generally lubricate the rubbing contacts only at high temperatures due to the formation of a viscous silicate melt layer. However, silicates behave such as rigid solid at low temperatures, where the friction is almost unaffected by the deformation strain rate. Lubrication is quite essential to improve the energy efficiency and durability of tools. The presence of inorganic sodium metasilicate at 920 °C for high-temperature friction tests of steel-to-steel contacts can reduce the friction coefficient to half and minimize one order of magnitude of wear rate [170].

Wang et al. [171] developed a magnesium silicate hydroxide-MoS$_2$-Sb$_2$O$_3$ nanocomposite coating for the high-temperature super-lubricity application. This composite coating with a thickness of 150–250 nm is accomplished by uniformly burnishing hydrothermally synthesized lamellate magnesium silicate hydroxide, molybdenum disulfite, and antimony trioxide powders onto a copper substrate. With increasing the temperature up to 400 °C, the friction coefficient of the composite coating rapidly decreases to below 0.01 within 100 revolutions and finally reaches a super-lubricity state, which is attributed to the synergistic characteristics of lubricating antimony trioxide, molybdenum disulfide,
and magnesium silicate hydroxide phases, and enables easy shearing. Magnesium silicate hydroxide is a type of layered silicate that releases unsaturated Si-O-Si, O-Si-O, OH-Mg-OH, OH-, and O-H-O groups, thus forming a tribofilm during sliding [172]. The magnesium silicate hydroxide-C-Sb2O3 coating burnished on a nickel superalloy substrate also exhibits high-temperature super-lubricity due to the formation of an easy-to-shear silicate-containing carbon film tribochemically-activated during sliding [172].

4.9. Cesium Oxythiomolybdate Cs2MoOS3

Research on cesium oxythiomolybdate (Cs2MoOS3), zinc oxythiomolybdate (ZnMoOS3), and cesium oxytrithiotungstate (Cs2WOS3) was initiated as a response to the requirement for a high-temperature solid lubricant for ceramic bearings in lightweight single-use engines. The US Air Force Research Laboratory at Wright-Patterson Air Force Base (WPAFB) developed a thin bonded lubricious coating of the complex chalcogenide, Cs2MoOS3, which could adequately lubricate silicon nitride bearings in the air at temperatures up to 760 °C, speeds up to 1.2 million DN (DN = bearing bore size in mm × shaft rpm) and thrust loads of 890 N for periods of five to six hours in 1987.

Cesium oxythiomolybdate-based lubricants have been applied by burnishing; however, the burnished powders need to often be replenished to prevent failure and ensure a long wear life. Pulsed laser deposition is an excellent method for highly adherent Cs2MoOS3 films due to its characteristic of replicating the target chemistry in the film. Therefore, Cs2MoOS3 films were developed to react with the environment to form new lubricious phases at elevated temperatures for the lubrication of silicon nitride bearings by pulsed laser deposition [173,174]. Figure 9 shows the friction behavior of Cs2MoOS3 films pulsed laser deposited on Si3N4 in sliding against Si3N4 ball [174]. This adaptive lubricant provides a low-friction surface with an average friction coefficient value of 0.03 at 600 °C. Not only that, room temperature and 700 °C are suitable as well; however, it does not work at temperatures of 300 °C and especially 800 °C. A high-temperature lubricious phase is formed based on the reaction of the components with each other and O2 with the increase in test temperature. Cesium oxythiomolybdate (Cs2MoOS3) used in a coating with a sodium silicate binder exhibited a friction coefficient of below 0.2 at 650 °C [174]. However, one challenge in using Cs2MoOS3 as a lubricant is its thermal instability above 200 °C. Oxidation products of Cs2MoOS3 powders at 600–800 °C consist of Cs2SO4, cesium oxides, Cs2MoO4, and other molybdenum oxides. Oxidation of Cs2MoOS3 to form lubricious Cs2MoO4, MoO3, and other oxides on the sliding surface provides good lubrication on ceramics at temperatures between 300 and 600 °C. Cs2MoOS3 films on the substrates of alumina and zirconia have friction coefficients of below 0.2 when sliding against an alumina ball. However, Cs2MoOS3 films lubricate well on Si3N4 substrates between 600 and 750 °C, and on SiC substrates between 500° and 600 °C, with a friction coefficient of below 0.1 in sliding against silicon nitride ball. The lubrication mechanisms depend mainly on the oxide softening and the formation of cesium silicate glass on the sliding surface [174].

Cesium oxytrithiotungstate (Cs2WOS3) and zinc oxythiomolybdate (ZnMoOS3) are also thermodynamically stable and have exceptional potential for lubrication of silicon nitride bearings at elevated temperatures [6,174]. The formation of a low shear strength glass was previously postulated as the lubrication mechanism for cesium tungsten (Cs2WO3) bonded coatings (containing lubricious cesium silicate glass) on silicon nitride bearings by Rosado et al. [175]. The high-temperature rolling contact endurance and tribological properties of lubricious cesium-based compounds were investigated on silicon nitride balls at 650 °C when applied as either a bonded or fused coating [6]. The best comprehensive performance was achieved with a cesium silicate reaction film formed in situ by Cs2O•SiO2 and a hydrated cesium silicate bonded coating of Cs2O•3SiO2•nH2O, which provided rolling friction coefficients less than 10−3 and small wear coefficients to obtain very long endurance lives at high contact stresses. In fact, alkali ions are known to reduce the viscosity of silicate glasses, which have been used as lubricants for steel working for over 70 years. Generally, glass lubricants with desirable properties above 600 °C can lubricate...
the hot extruded steel surfaces at large area reductions and in longer lengths [176]. Since thin PLD films are not suited to long-term use, they are most advantageously employed for short-duration and one-time applications. Experimental prototypes of other engine parts (e.g., piston pins, intake and exhaust valves, camshafts, cam lobes, roller followers, high-temperature seals, etc.) made from Si$_3$N$_4$ must be considered to further evaluate their tribological performance and reliability.

Figure 9. Friction behavior of Cs$_2$MoOS$_3$ films pulsed laser deposited on Si$_3$N$_4$ in sliding against Si$_3$N$_4$ ball. Reproduced with permission from Reference [174], Copyright © Elsevier Science B.V. 2002.

4.10. Synergistic Effects of Various Solid Lubricants

Lubrication problems are particularly challenging in the aerospace industry since aerospace components experience high and low temperatures, alternative moist air and vacuum, heavy load, high speed, and strong radiation conditions [177]. For example, lubricants that are effective over an extreme temperature range are necessary for new-generation advanced engines with increased propulsion. Lubricants for space vehicles must be able to function in moist air through a vacuum, such as on reusable launch vehicles. The coated satellite components designed for long lifetimes may be tested and stored in a terrestrial environment before they are operated in the vacuum of space. Unfortunately, it is particularly difficult to find a single solid lubricant that can provide the best lubrication performance over all conditions. As no individual solid lubricant material can provide adequate lubrication properties under environmental conditions of low/high temperature, high radiation, high pressure, high chemical reactivity, and ultrahigh vacuum, a microstructurally engineered combination of various solid lubricants will become necessary for advanced lubrication systems [177].

Before an optimized solid lubricant combination is microstructurally designed, it is of great importance to understand the fundamental characteristics and fabricating methods of different constituents. Soft metals of Ag and Au contain multiple slip planes and produce enhanced ductility and easy-to-shear tribolayer. Graphite, h-BN, and MoS$_2$ are typical layered materials with easy-to-shear interlayers. Nb(S, Se, Te)$_2$ is used in applications requiring good electrical conductivity. CaF$_2$ or BaF$_2$/CaF$_2$ are common high-temperature lubricants (i.e., >500 °C), while PbO has good lubricating properties from about 450 to 700 °C, although it is still limited by its low temperature shear strength and by phase transition, reactivity, and melting at high temperatures. Figure 10 shows effective operating temperature ranges for a variety of solid-lubricating materials [20]. The synergistic effects of different solid lubricants have been widely explored for humidity-, temperature-, vacuum, or load-adaptive tribological applications.
Typical examples associated with these combinations of various solid lubricants are quite inspiring for specific engineering applications such as aircraft and satellites, where continuous variations from humid air to vacuum during launch and flight or return periods over a broad vacuum range. Graphite, DLC, and MoS$_2$/WS$_2$ powders are combined to lubricate across humid air (graphite, DLC) and vacuum (MoS$_2$/WS$_2$). Multiple mechanisms for air-space humidity-adaptive response of W-C-S, W-C-Se, YSZ-Au-MoS$_2$-DLC, and Al$_2$O$_3$-Au-MoS$_2$-DLC nanocomposite coatings are designed for lubrication subjected to environmental cycling [3]. Interestingly, the 500 nm thick W-C-S coating exhibited a friction coefficient comparable to that of pure graphite in humid air and simultaneously attained the equivalent friction coefficient of pure WS$_2$ in vacuum for more than $2 \times 10^6$ cycles with a wear rate less than $10^{-7}$ mm$^3$/(Nm). The sensitivity of MoS$_2$ and WS$_2$ coatings to relative humidity can be reduced through the use of dopants, additives, or interlayers such as W-C-S and W-C-Se coatings. A novel scheme to operate across a range of climatic conditions was described whereby a circular groove was laser milled into a TiC/DLC composite coating and filled with MoS$_2$ additives. MoS$_2$ additives form a transfer film and generate lubrication in a vacuum and dry N$_2$, while a graphite-like transfer film provides lubrication in moist air. A promising path of incorporating multiple lubricants in a composite coating to reduce friction and wear will emerge for a useful number of cycles through a broad range of ambient humidity and temperatures on a reusable air-space vehicle capable of multiple atmospheric reentries.

Sliney and Dellacorte pioneered self-lubricating composites or coatings comprising silver for low-temperature lubrication, BaF$_2$-CaF$_2$ eutectic for high temperatures, and Cr$_3$C$_2$ or Cr$_2$O$_3$ for enhanced wear resistance in tribological applications such as foil air bearings and mechanical seals of advanced propulsion systems. This lubrication combination of silver-BaF$_2$/CaF$_2$ eutectic is quite thermally stable over a wide temperature range, which can be cycled to provide self-lubricity between low and high temperatures for long-term service [1,7,93,94]. Similarly, the addition of both gold and CaF$_2$ into ZrO$_2$(Y$_2$O$_3$) ceramics significantly reduces the friction and wear up to 800 °C tested with a ball-on-disc configuration. Interestingly, gold was not able to entirely cover the contact region at room temperature by diffusion, but it does reduce friction and wear significantly [102]. The enhanced wear resistance is attributed to the lubrication of discontinuous protective layers at moderate temperatures or nano-scale gold particles on CaF$_2$ layers at high temperatures.
The use of gold as an effective solid lubricant is specialized for lubrication applications in aerospace tribology or the protection of electronic connectors [2].

Typical examples of synergistic lubrication in MoS$_2$-containing composite coatings include PbO-MoS$_2$ [117,178], MoS$_2$-Sb$_2$O$_3$, YSZ-Ag-Mo-MoS$_2$ [179], YSZ-MoS$_2$-DLC-Au [3], Mo$_2$N-MoS$_2$-Ag [119], Ni-Al-Ag-MoS$_2$ and Ni-Al-Ag-MoS$_2$-hBN [180]. These coatings provide effective lubrication protection over a broad temperature range in a combination of the lubricity of MoS$_2$ at lower temperatures and vacuum with the in situ formation of easy-to-shear lubricious oxides at elevated temperatures [30,181,182]. However, adaptive lubrication surfaces by Au or Ag may have some challenges, including the random migration of soft metal clusters, poor adhesion in the presence of oxygen, and excessive deformation at elevated temperatures on the sliding surface. It is particularly important to note that the tribo-chemical reactions to the lubricious film are not reversible, and the tribo-chemical induced oxides do not provide lubrication at low temperatures for repeated thermal cycling [8]. The primary adaptive mechanisms are associated with environmental-assisted oxidation to form easy-to-shear and low-melting-point binary TiO$_2$, V$_2$O$_5$, MoO$_3$ Magneli phases and ternary oxides including silver molybdates, vanadates, tungstates, niobates, and tantalates, etc., temperature-activated diffusion of soft metals, and thermo-mechanically induced phase transitions to reorient hexagonal solids and promote surface-hardening [30,31]. Adaptive lubricants must be engineered due to the irreversible tribo-chemical reactions so that they can be cycled between low and high temperatures [177].

The challenge associated with this issue was the reversibility of the thermally adaptive tribological surfaces over multiple thermal cycles, analogous to their humidity- or vacuum-adaptive counterparts occurring in various engineering applications. Several approaches are postulated for adaptive multilayered coatings and surface multifunctional design, such as bionic compositing, tunable surface texturing, and tribo-reaction of oriented lubricants. For example, temperature-adaptive composites exhibiting diffusion-, melting-, or oxidation-limiting lubrication are developed over multiple thermal cycling through microlaminate architectures in engineered designs to activate the functionality of various solid lubricants [30,177]. A microlaminate coating comprising alternative adaptive coating layers and diffusion barrier layers would inhibit the diffusion and migration of Ag or Au clusters from the underlying layers and restrict the diffusion of oxygen into the coating to generate tribo-oxidation until needed [3]. Thus, both the diffusion and migration of soft metallic clusters and the formation of lubricious oxide would occur on demand in such a microlaminate coating.

5. Production Technologies of Self-Lubricating Composites and Coatings

There are different approaches for applying high-temperature solid lubricants for a variety of engineering applications [21]. (1) Powdered solid lubricants have obvious shortcomings, in spite of powder lubrication as the oldest and simplest method. Two factors impeding the development of powder lubrication are associated with the difficulty of forming a continuous solid lubricating film and the need to constantly replenish the lubricant during the wear process. (2) Bonded coatings supply a larger film thickness and an enhanced wear lifetime, which improves the reliability and durability of traditional solid lubrication. However, most resin- or inorganic-bonded coatings lubricate only at low temperatures and are not suitable for high-temperature conditions. (3) Surface texturing of sliding metallic [183,184] or ceramic components/coatings [185] by laser [186], pulsed electric arc [187], ion etching, and electron beam followed by filling of various solid lubricants such as graphite, MoS$_2$/WS$_2$ [188], Mo/MoS$_2$/Ag [51], CaF$_2$, etc. (4) Self-lubricating composites or coatings are the most widely used and most promising solid lubrication technologies. A variety of processing approaches of large-scale two-dimensional layered films from graphene, h-BN, and TMD, to black phosphorous nanosheets and MXenes for solid lubrication as shearing films and nano-roller bearings, have been developed rapidly with the advent of graphene, which includes mechanical or chemical exfoliation, atomic/molecular
layer deposition, chemical vapor deposition, magnetron sputtering, pulsed laser deposition, electrophoretic deposition, ink-jet printing, spray/spin coating, or directly burnishing onto the substrate. Preparation techniques of self-lubricating composites or coatings include physical/chemical vapor deposition, thermal/plasma spraying, electrodeposition, laser cladding, powder metallurgy processes such as pressureless sintering, hot pressing, hot isostatic pressing and spark plasma sintering, and hot rolling.

5.1. Self-Lubricating Coatings by Physical Vapor Deposition

Physical vapor deposition processes are versatile in the composition of coating materials. They have the ability to produce very thin coatings as low as several nanometers with high purity, high adhesion, and unusual microstructures at high deposition rates. Physical vapor-deposited coatings for friction and wear applications include the following two different types: soft solid-lubricant coatings and hard wear-resistant coatings. Applying PVD coatings includes simple vapor deposition, sputtering, ion plating, and pulsed laser deposition.

Recently, various ion beam-assisted deposition methods, referred to as modifications of the more basic process, have been used to deposit very adherent tribological coatings at relatively low deposition temperatures. Strongly adherent MoS$_2$-Au(Ag)-Re composite films deposited by PVD combined with ion bombardment exhibit columnar structure and five times higher wear resistance than conventional sputtered MoS$_2$ film. Besides the sputtered MoS$_2$ films, two other categories of PVD coatings that are tribologically significant are sputtered hard coatings for wear control, such as diamond-like carbon coatings doped with or without additives, and ion-plated soft metals (Au, Ag, or composites) for lubrication, such as magnetron sputtering assisted pulsed laser deposition (MSPLD). Adaptive WC-WS$_2$-DLC, YSZ-Au-DLC-MoS$_2$ coatings are designed to provide graphite-like lubrication in humid air and molybdenum and tungsten sulfite lubrication in dry sliding and vacuum [3]. Gradient multilayered soft metal films of Ni-Cu-Ag-In significantly improve the load capacity and mechanical properties of films on space harmonic gears and rocket turbo-pump bearings. Sputtering is probably the most widely used vapor deposition technique. Nano-crystallization drove high-temperature self-lubricating WS$_2$ coatings deposited by magnetron sputtering, which have a low friction coefficient of 0.07–0.2 at temperatures of 100–400 °C. At 500 °C, the oxidation-induced formation of WO$_3$ with a relatively low hardness reduces the friction and improves wear resistance [82]. A self-lubricating Al$_2$Au-containing Al-Au coating was deposited onto cemented carbide substrates using unbalanced magnetron sputtering, which exhibits a fine-grained structure and a hardness value of 4 GPa. The coatings are thermally stable up to about 850 °C, and then the onset of oxidation occurs, and tribo-oxidation contributes to a low friction coefficient when testing against alumina tribopairs at temperatures of 500 °C to 700 °C [189]. RF magnetron sputtering was used to deposit periodic CrAlN/VN multilayer coatings with various bilayer periods, which provide preferable anti-wear performance due to interfacial coherent strengthening and lubricious vanadium oxide layers formed on the sliding surface [190]. In reactive sputtering, a reactive gas is introduced to produce compound coatings. Silver tantalate coatings of the Ag-Ta-O system produced by unbalanced magnetron sputtering exhibit excellent tribological properties at high temperatures [191]. Self-lubricating ceramic-Ag coatings were investigated due to the large variety of functionalities (e.g., lubricity, surface plasmon resonance, and antibacterial effect) that silver possesses and the possibility of selecting the proper matrix to support the metallic lubricant [192].

Researchers at the USA Air Force Research Laboratory, Wright-Patterson Air Force Base (WPAFB), developed PVD deposition methods for lubricous adaptive coatings, where initial multiphase adaptive oxide and fluoride coatings for high-temperature lubrication were progressively evolved to hard oxide matrices with additions of easy-to-shear gold and silver for moderate temperatures and dichalcogenide and graphite for near room temperature lubrication [3,30]. Thin composite films of MoS$_2$ with PbO, WS$_2$ with ZnO, MoS$_2$ with Sb$_2$O$_3$, and later WS$_2$ with CaF$_2$ are deposited to produce lubrication over
a wide temperature range. In this case, the metal dichalcogenides reacted with the oxides to form lubricious PbMoO$_4$, ZnWO$_4$, and CaSO$_4$ phases at elevated temperatures [126,127]. However, the drawback to this concept is that the reaction is irreversible, and the low-temperature lubrication will lose inevitably once returning to room temperature during the thermal cycling process.

5.2. Self-Lubricating Coatings by Thermal Spraying

The thermal spraying process has nowadays become a cost-efficient and reliable method to deposit thick coatings with a wide variety of feedstock materials and substrates, which has been successfully applied in aerospace, aviation, and power generation, becoming an essential component of modern industry [193]. The thermal energy necessary to melt the spraying material in the form of powder, wire, or even liquid suspension can be produced by a flame created by combustion gases, the electric plasma produced by an electric discharge, the detonation of the combustion gas by a spark plug, or an electric arc. Generally, only those thermally stable feedstock materials without the occurrence of oxidation or decomposition can be sprayed. In a plasma spraying process, the spraying feedstocks are fed into the plasma flame by a carrier gas, where they become melted or semi-melted and are then propelled to the substrate at high velocity due to high plasma enthalpies.

Plasma-sprayed ceramic and cermet coatings, especially those based on oxides (Al$_2$O$_3$, Cr$_2$O$_3$, etc.) and carbides (WC, Cr$_3$C$_2$, etc.), are widely used for wear and corrosion protection. Solid lubricants of graphite, MoS$_2$, fluorides, and lubricious binary or ternary oxides embedded in plasma sprayed coatings have been employed in different industries, especially in those applications under severe operation conditions [194,195]. The status-of-the-art developments involve reactive plasma spraying and plasma-sprayed oxide ceramic coatings containing nanophases either from nanostructured precursors or post-precipitated by purposely designed thermal treatments [180].

Composite Magneli phases of Ti$_6$Cr$_2$O$_7$ and CrTi$_2$O$_5$ from the system Ti$_n$−$2$Cr$_2$O$_{2n−1}$ formed on the plasma-sprayed TiO$_2$−Cr$_2$O$_3$ coating provide solid lubrication to 800 °C during wear tests [108]. Ni-Al-Cr$_2$O$_3$−Mo−Ag coatings were plasma sprayed on the surfaces of Inconel 718 alloy. The sliding process at elevated temperatures could promote the formation of lubricious silver molybdate films to reduce friction and wear [196]. Plasma sprayed WC-Co-Cu-BaF$_2$/CaF$_2$ self-lubricating coatings not only offer effective solid lubrication but also self-heal the surface defects and protect WC from decarburization [195]. The optimized composite coating consists of 70 wt.% WC-(W,Cr)$_2$C-Ni, 15 wt.% Ag, and 15 wt.% BaF$_2$-CaF$_2$ exhibits excellent friction and wear performance over a wide range of temperature up to 600 °C [197]. The synergistic effects of Ag, MoS$_2$, and hBN help to improve the tribological performance of Ni-Al-Ag-MoS$_2$ and Ni-Al-Ag-MoS$_2$-hBN on the surface of titanium- and nickel-matrix alloys [180].

Researchers from the NASA Glenn research center developed different self-lubricating plasma-sprayed coatings, such as PS100, PS200, PS300, PS304, and PS400, for extreme temperature applications. The objective of these composite coatings is to incorporate these fluorides and soft metals (i.e., Ag and Au) into matrix materials to reduce friction and wear at elevated temperatures. PS100 was a plasma sprayed nickel-chromium-based coating containing glass as a binder and silver and fluorides as the solid lubricants for applications of low speed, oscillating, plain spherical bearings. PS100 exhibited a very low friction coefficient and relatively low wear resistance over a broad range of temperatures, which was designed for high-temperature applications involving moderately loaded, up to at least 34 MPa, sliding contact bearings and for shaft seals requiring a degree of conformability [1]. PS200 was a plasma-sprayed nickel-cobalt-based chromium carbide coating, with silver and BaF$_2$-CaF$_2$ eutectic fluorides as the solid lubricants. This provides better tribological performance at temperatures as high as 900 °C for applications in the cylinder walls of Stirling engines. A linear alternator is a reciprocating device that is driven by a solar or nuclear-powered Stirling engine to generate electrical power. The power piston and cylinder were made of Ti-6Al-4V alloy and were designed to be lubricated by
a hydrodynamically-generated gas film, where rubbing was generated during starts and stops and there was the possibility of an occasional high-speed rub to lead to a severe galling tendency in sliding contacts, and thus a back-up self-lubricating coating on the cylinder and/or piston was needed [93,95,198]. PS212 coatings could be used for foil gas bearing applications [194]. PS300 was a plasma sprayed nickel-chromium bonded chromium oxide coating with 60 wt.% Cr2O3, 20 wt.% Ni80Cr20, 10 wt.% silver, and 10 wt.% BaF2-CaF2 eutectic solid lubricant additions, which provided good friction and wear properties in a partial-arc foil bearing at 500 °C. The PS300 coating was successful in lubricating a foil gas bearing at 500 °C for over 15,000 start-stop cycles. However, excessive foil wear was still observed at room temperature, and journal coating adhesion was poor, especially during repeated thermal cycles, which led to the spalling of PS300 coating (delamination) directly from the journal surface due to a mismatch in thermal expansion coefficients. Subsequent research resulting in a modification to the PS304 coating could achieve low friction and a small wear rate at elevated temperatures up to 650 °C [94]. PS304 is a 60 wt.% Ni80Cr20 matrix that contains solid lubricants, 10 wt.% Ag and 10 wt.% BaF2-CaF2 in eutectic composition. In total, 20 wt.% chromium oxide particles were used for strength enhancement, while Ag and BaF2-CaF2 were used to provide lubrication properties at different temperatures [199]. PS304 was found to be a suitable coating for many applications of foil air bearings and seals, exhaust gas recirculation (EGR) and turbocharger waste gate valve stems, steam turbine control valve components and bushings and shafting in heat treatment furnaces [200]. PS304 coating exhibits an increase in adhesive strength after exposure to temperatures above 500 °C for various durations due to the precipitation of a chromium-rich second phase in the matrix. Heat treatment of PS304 coating after plasma spraying and prior to finish machining is needed to insure dimensional stability and maximum adhesive properties [201]. One example is the development of a self-lubricating PS304 coating on advanced high-temperature nickel-based superalloy foil air bearings to evaluate their performance and durability from 25 °C to 650 °C [202,203]. In order to achieve the PS304 surface smoothening effect that is noted in high-temperature air foil bearings subject to cyclic start/stop operation, thrust-washer tests were also conducted with intermittent, as opposed to continuous, sliding contact [204,205]. The foil surface experiences sliding contact with the shaft during the initial start/stop operation prior to developing a gas film, which may lead to wear and failure due to inadequate load capacity, material capabilities, or rotordynamic performance. These tested bearings with PS304 coatings provided lives well in excess of 30,000 cycles, particularly some of them even exceeding 100,000 cycles, and are well suited for advanced high-temperature oil-free turbomachinery applications. Another example involves the development of self-lubricating steam turbine lift rods, which are designed to work in a frictional contact mode against a stack of seal rings at elevated temperatures and face challenging needs for surface protection. Wang et al. [206] reported a high-temperature self-lubricating composite coating deposited by plasma spraying to overcome the wear and galling damage of steam turbine governor valve lift rods of Waspaloy, a nickel-based alloy (AMS 5604), subjected to metal-metal interaction at 540 °C. A PS304 coating applied on the lift rods was intact even after 8500h of operation at 540 °C. Field operation after 3 years up to the date indicated that PS304 coatings have functioned effectively. A lubricious glaze layer containing Cr2O3, Ag, and CaF2/BaF2 eutectic was formed on the surface of coated lift rods to effectively protect them from high-temperature wear and galling damage. PS400 was a plasma-sprayed nickel-molybdenum-aluminum matrix containing chromium oxide as a binder, while the silver acted as a solid lubricant at a relatively lower temperature range of room temperature to approximately 450 °C, and BaF2-CaF2 eutectic offers effective lubricating above 400 °C, respectively. Therefore, PS400 coating was excellent for high-temperature wear applications due to the formation of a self-lubricating glaze layer, and it was used for microturbine engine foil gas bearing applications at temperatures above 760 °C [207]. The PS400 coating was tested as a shaft coating operating against a hot section foil bearing in a Capstone oil-free 30 kW microturbine engine, and interestingly, the shaft
diameter measurements show almost no wear due to the development of lubricious surface glazes after 2200 start-up and shut-down cycles conducted over 8000 h of operation at 96,000 rpm and approximate 540 °C. These coatings developed by NASA Glenn Research Center were effective at elevated temperatures but typically had friction coefficients above 0.2 at room temperature. In addition, the plasma spray technique intrinsically produces very thick (typically hundreds of microns) coatings, limiting their utility in application to precision components.

Plasma-sprayed ZrO$_2$-CaF$_2$ coatings incorporated with and without silver addition exhibited attractive tribological properties in sliding against superalloys [1]. Both coating combinations had fairly high wear rates at room temperature, but wear rates were much lower for the ZrO$_2$-CaF$_2$ coating at 650 °C. The effect of different lubricants on the high-temperature friction and wear characteristics of low-pressure plasma-sprayed ZrO$_2$(Y$_2$O$_3$)-based ceramic coatings was investigated [15,16,208]. Low friction and wear at temperatures of 600–700 °C were achieved for the low-pressure plasma-sprayed ZrO$_2$-CaF$_2$ coating when sliding against an alumina ball [16]. The incorporation of CaF$_2$/Ag$_2$O lubricants into the self-lubricating ZrO$_2$(Y$_2$O$_3$) matrix composite coating provided low friction and wear against an alumina ball at temperatures of 300 °C to 700 °C [208]. However, high friction and wear of coatings were observed due to excessive plastic deformation and material transfer at 800 °C. Low-pressure plasma sprayed ZrO$_2$(Y$_2$O$_3$) matrix coating incorporated with 50 vol.% BaCrO$_4$ ternary oxide effectively reduced friction and wear at temperatures of 400 °C to 800 °C on reciprocating ball-on-flat tribometer due to the formation of easy-to-shear BaCrO$_4$-rich tribolayers [15]. The incorporation of CaF$_2$, BaF$_2$, and Cr$_2$O$_3$ into a plasma-sprayed composite coating also achieved friction coefficients of 0.20–0.30 from 300 °C to 900 °C, which was due to the coexistence of CaF$_2$ and BaCrO$_4$ induced by the tribo-chemical reaction on the sliding surfaces. Chromium oxide coatings are widely used for tribological applications requiring high wear resistance and lubricity at elevated temperatures [17]. The incorporation of BaCrO$_4$ into chromium oxide coatings was developed to further improve its tribological properties as a high-temperature solid lubricant up to 700 °C. The adaptive formation of lubricious chromate films on worn surfaces of plasma-sprayed Cr$_2$O$_3$-BaCrO$_4$ coatings plays a vital role in reducing friction and wear at elevated temperatures [209].

Spray-formed and powder metallurgy materials are usually of high porosity. As one of the secondary processing techniques, hot rolling or forging is used to densify the products and improve their properties [210]. The Al-Si-graphite self-lubricating composite was synthesized by the spray forming process and then densified by warm rolling. Tribological tests were performed in the pin-on-disk method with high-carbon EN 31 steels as the counter surface at room temperature. Finally, the friction coefficient and wear rate were reduced from 0.49 to 0.38 and 2.25 × 10$^{-3}$ mm$^3$/(Nm) to 1.25 × 10$^{-3}$ mm$^3$/(Nm), respectively, with the thickness reduction of the composite by warm rolling [210].

5.3. Self-Lubricating Composites by Powder Metallurgy

Generally, wear-resistant components with flat surfaces and outside diameter surfaces are readily accessible for a plasma spraying process. However, the inside surfaces of relatively small components such as small cylindrical bearings, bushings, and valve guides are difficult or impossible to deposit coatings from a plasma spray gun. Powder metallurgy processes are appropriate for these types of components. During sintering, various mass transfer processes contribute to the densification of raw powders consisting of matrix materials and solid lubricants. Figure 11 shows the typical microstructure and worn surfaces of a spark-plasma-sintered ZrO$_2$(Y$_2$O$_3$)-30CaF$_2$-30Au composite. The obtained self-lubricating materials usually exhibit uniformly distributed grains due to the growth inhibition effect of different constituents [102]. The hot-pressed alumina matrix composite incorporated with Ag and CaF$_2$ provided low friction and wear at temperatures of 300–750 °C [211–213]. The appropriate number of solid lubricants and sintering aid
appears to improve the formation of a complete, well-adhering self-lubricating film, which seems to be responsible for the reduction of friction and wear at 650 °C.

\[\text{Figure 11. Microstructure and worn surfaces of a ZrO}_2(\text{Y}_2\text{O}_3)-30\text{CaF}_2-30\text{Au composite fabricated by spark plasma sintering: (a) Morphology and phase distribution of different constituents, (b) worn surface after room temperature wear test, (c) worn surface after 400 °C wear test, (d) worn surface after 800 °C wear test. Reproduced with permission from Reference [102], Copyright © Elsevier B.V. 2004.}\]

\(\text{ZrO}_2(\text{Y}_2\text{O}_3)-50\text{BaCrO}_4\) composites were spark plasma sintered by optimizing the composition and sintering parameters, and high-temperature friction and wear tester was used to evaluate their friction and wear properties in dry sliding against an alumina ball. The sintered samples exhibit a density of 4.99 g/cm\(^3\) and a Vickers hardness of 426 HV\(_1\) and have coefficients of friction from 0.38 to 0.55 and wear rates from \(1.44 \times 10^{-5}\) to \(5.35 \times 10^{-5}\) mm\(^3\)/Nm from room temperature to 800 °C [143]. A well-covered BaCrO\(_4\) fine-grain layer is observed on the sliding surfaces at elevated temperatures due to considerable plastic deformation and smearing of BaCrO\(_4\). The synergistic effects of CaF\(_2\) and Au lubricants on the friction and wear properties of ZrO\(_2(\text{Y}_2\text{O}_3)\) matrix composites were evaluated with a high-temperature friction and wear tester. The ZrO\(_2(\text{Y}_2\text{O}_3)\)-CaF\(_2\)-Au composite has a density of 5.53 g/cm\(^3\) and a Vickers hardness of 396 HV\(_1\) and exhibits a friction coefficient from 0.36 to 0.50 and a wear rate from \(1.67 \times 10^{-6}\) to \(3.55 \times 10^{-6}\) mm\(^3\)/Nm over the entire temperature range up to 800 °C [102]. Soft metallic lubricants of gold are observed to diffuse and migrate from the subsurface layer to the sliding surface and finally form a discontinuous lubrication film, which lubricates the oxide matrix synergistically with CaF\(_2\). From an engineering standpoint, most machinery runs at low temperatures only during start-up periods; therefore, low friction (<0.30) at high operating temperatures, especially above 300 °C, for a gas turbine seal, for example, is of significant value.

Hot pressing has been used to fabricate NiCr-BaCr\(_2\)O\(_4\) self-lubricating composites in a vacuum sintering furnace for tribological evaluation in sliding against an alumina ball. At 800 °C, a friction coefficient of 0.27 and a wear rate of \(4.5 \times 10^{-6}\) mm\(^3\)/Nm are achieved for the NiCr-20BaCr\(_2\)O\(_4\) composite, however, for a comparative study, the unmodified NiCr alloy exhibits a friction coefficient of 0.41 and a wear rate of \(9.2 \times 10^{-5}\) mm\(^3\)/Nm [151]. A dense oxide glaze layer is formed on the sliding surface of the NiCr-20BaCr\(_2\)O\(_4\) composite.
to effectively reduce friction and wear at elevated temperatures, and the tribo-oxidation reaction of BaCr$_2$O$_4$ is responsible for the formation of the glaze layer containing lubricious hexavalent-Cr$^{6+}$ BaCrO$_4$ during high-temperature wear tests.

Self-lubricating ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$-Ba$_2$Sr$_{1-x}$SO$_4$ ($x = 0.25, 0.5, 0.75$) composites were fabricated by spark plasma sintering. The composites containing Ba$_2$Sr$_{1-x}$SO$_4$ have significant improvements in effectively reducing friction and wear in sliding against an alumina ball as compared to unmodified ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$ ceramics. Interestingly, the composites have low friction coefficients of less than 0.15 and wear rates in the order of 10$^{-6}$ mm$^3$/(Nm) at 760 °C. For example, ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$-50Ba$_2$O$_2$SrO$_2$SO$_4$ composite has a density of 4.2 g/cm$^3$ and a hardness of 3.48 GPa, exhibits friction coefficients and wear rates of 0.2 and 4.8 × 10$^{-5}$ mm$^3$/(Nm) at room temperature, and 0.11 and 3.15 × 10$^{-6}$ mm$^3$/(Nm) at 760 °C, respectively. While ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$-50Ba$_{0.75}$Sn$_{0.25}$SO$_4$ composite has a density of 4.45 g/cm$^3$ and a hardness of 3.48 GPa, exhibits friction coefficients and wear rates of 0.24 and 2.06 × 10$^{-5}$ mm$^3$/(Nm) at room temperature, and 0.12 and 5.76 × 10$^{-6}$ mm$^3$/(Nm) at 760 °C, respectively. The formation and effective spread of the lubricious Ba$_2$Sr$_{1-x}$SO$_4$ layer is of great significance in lowering friction and wear [157]. Fe$_3$Al matrix composites containing Ba$_{0.25}$Sr$_{0.75}$SO$_4$ fabricated by hot pressing also exhibit a very low friction coefficient from 0.19 to 0.29 and wear rate on the order of 10$^{-5}$ mm$^3$/(Nm) at temperatures of 600–800 °C, which is lower than unmodified Fe$_3$Al alloy [158].

For a comparative study, a variety of self-lubricating ZrO$_2$(Y$_2$O$_3$) matrix composites are spark plasma sintered by tailoring the chemical compositions and sintering parameters. Selected lubricants of BaF$_2$, CaF$_2$, Ag, Ag$_2$O, Cu$_2$O, BaCrO$_4$, BaSO$_4$, SrSO$_4$, and CaSiO$_3$ were added to the oxide ceramic matrix to evaluate their self-lubricity over a wide temperature range up to 800 °C by using a ball-on-block friction and wear tester in sliding against alumina ball [101]. Figure 12 shows the frictional behavior of ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$ ceramics as a function of wear cycles at different temperatures [101]. Figure 13 shows the frictional behavior of ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$-50SrSO$_4$ composites as a function of the wear cycle at different temperatures [101]. The incorporation of graphite or Mo$_2$I$_2$ into ZrO$_2$(Y$_2$O$_3$) matrix composites was also evaluated for comparison under identical test conditions. The ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$ ceramics have a hardness of 14.8 GPa at room temperature and 7.8 GPa at 700 °C, and exhibit friction coefficients and wear rates of 0.48 and 2.31 × 10$^{-6}$ mm$^3$/(Nm) at room temperature, and 1.15 and 4.77 × 10$^{-4}$ mm$^3$/(Nm) at 800 °C, respectively. While, the ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$-50SrSO$_4$ composites have a density of 4.05 g/cm$^3$ and a hardness of 3.48 GPa, exhibits friction coefficients and wear rates of 0.11 and 1.05 × 10$^{-5}$ mm$^3$/(Nm) at room temperature, and 0.19 and 2.28 × 10$^{-6}$ mm$^3$/(Nm) at 800 °C, respectively. The dominating wear mechanism of unmodified ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$ ceramics depends mainly upon surface fatigue and brittle fracture at 800 °C. However, for self-lubricating composites, the most important factor for reducing friction and wear at elevated temperatures is attributed to the formation, plastic deformation, and effective spread of sulfate lubricating films on the sliding surface [101].

Figure 14 shows the frictional behavior of ZrO$_2$(Y$_2$O$_3$) matrix composites incorporated with and without various additives as a function of temperature in sliding against an alumina ball. Figure 15 shows the wear rates of ZrO$_2$(Y$_2$O$_3$) matrix composites incorporated with and without solid lubricants at room temperature and 800 °C [101]. Clearly, the incorporation of SrSO$_4$-CaSiO$_3$ solid lubricants into ZrO$_2$(Y$_2$O$_3$) matrix composites contributes to friction coefficients of 0.2–0.4 and wear rates of 10$^{-5}$–10$^{-6}$ mm$^3$/(Nm) from room temperature to 800 °C [101].

The synergistic effects of SrSO$_4$ and Ag on the self-lubricity of alumina matrix composites are evaluated by sliding against an alumina ball over a wide temperature range. The Al$_2$O$_3$-50SrSO$_4$-10Ag composites with a density of 3.69 g/cm$^3$ have a hardness of 129 HV$_1$, and exhibit friction coefficients and wear rates of 0.12 and 1.1 × 10$^{-5}$ mm$^3$/(Nm) at room temperature, and 0.22 and 4.89 × 10$^{-8}$ mm$^3$/(Nm) at 760 °C, respectively. While Al$_2$O$_3$-50SrSO$_4$-20 Ag composites with a density of 4.15 g/cm$^3$ have a hardness of 319 HV$_1$, and exhibit friction coefficients and wear rates of 0.08 and 1.06 × 10$^{-8}$ mm$^3$/(Nm) at room
temperature, and 0.20 and $3.57 \times 10^{-8}$ mm$^3$/Nm at 760 °C, respectively. The synergistic incorporation of SrSO$_4$ and Ag into the alumina matrix not only greatly improves friction and wear properties at low temperatures, but also provides excellent lubrication properties at elevated temperatures. From the friction and wear point of view, the incorporation of SrSO$_4$ and Ag into the alumina matrix creates a bright future for lubricating ceramic components in sliding or rolling contact over a broad temperature range up to 800 °C. The strontium sulfate lubricants become softened at high temperatures, and further spread out onto the sliding surface as well as the soft Ag clusters extrude and migrate from the subsurface layer due to tribological stressing, forming a lubricious mechanically mixed layer of SrSO$_4$ and Ag [163]. In previous work, the oxometallate-containing coatings were also fabricated by various powder metallurgy processes, such as Al$_2$O$_3$-BaCrO$_4$ coatings by hot pressing and SrSO$_4$-Sr$_2$SiO$_4$ coatings by spark plasma sintering on the surfaces of oxide ceramic components to improve the tribological properties.

Figure 12. Frictional behavior of ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$ ceramics as a function of wear cycles at different temperatures. Reproduced with permission from Reference [101], Copyright © Elsevier B.V. 2009.

Figure 13. Frictional behavior of ZrO$_2$(Y$_2$O$_3$)-Al$_2$O$_3$-50SrSO$_4$ composites as a function of wear cycle at different temperatures. Reproduced with permission from Reference [101], Copyright © Elsevier B.V. 2009.
Spark plasma sintering has also been used in the preparation of self-lubricating composites containing graphene. Hybrid Al-matrix composites containing nano-Al2O3 and graphene nanoplatelets were prepared in an argon atmosphere at 500 °C. The friction coefficient decreased from 0.36 to 0.24 against chromium-plated the chrome steel counter body by increasing the content of graphene nanoplatelets from 0.5 wt.% to 5 wt.% [214].
The NiAl matrix composites with 1.5 wt.% graphene nanoplatelets were fabricated at 1150 °C and then the tribological tests were performed by sliding against a Si$_3$N$_4$ ball. The friction coefficient reaches 0.36 at 200 °C. On the other hand, the wear rate increases rapidly with temperature, reaching $3.39 \times 10^{-4}$ mm$^3$/Nm at 400 °C, which may be attributed to surface layer delamination due to the stress-inducing crack propagation parallel to the sliding direction [215]. Moreover, the synergetic effects of multilayer graphene and Ti$_3$SiC$_2$ have been studied in TiAl and Ni$_3$Al matrix composites. Tribological evaluation at various temperatures shows that multilayer graphene enriched on the worn surface can effectively reduce friction and wear below 350–400 °C but lose effect at higher temperatures. Nevertheless, Ti$_3$SiC$_2$ will partially decompose and form a lubricating film on a worn surface at 400–800 °C, which contributes to the outstanding high-temperature tribological performance [216,217]. Friction stir processing was also used to obtain an aluminum-graphene composite. The friction tests were performed using the block-on-ring method on a worn surface at 400–800 °C, which contributes to the outstanding high-temperature tribological performance [216,217].

5.4. Self-Lubricating Coatings by Electrodeposition

Electrochemical deposition (also known as electrolytic deposition, electrodeposition, or electro-plating, electro-spark deposition) is the widely used and convenient technology of applying metallic coatings with relatively high melting points. Coatings with a thickness of sub-micro to tens of microns can be applied to a metallic surface (e.g., Ni, Co, Cu, Pb, Cr) and, with suitable preparation, to plastics and many other nonconductive substrates at room temperature or slightly higher temperatures of less than 100 °C. Codeposition of two or more metals is possible under suitable conditions of electric potential and polarization. A wide range of composite coatings containing insoluble particles in a metallic matrix can be fabricated by codepositing particles from suspension in agitated electrolytic solutions. The evergrowing demands of industrial surface engineering, such as wear- or corrosion-resistant surface layers, provide a driving force for rapid development of electrodeposition over the decades [219].

Electrodeposition of nickel-matrix composites containing dispersions of second-phase particles has attracted significant importance for potential applications. Incorporation of hard oxides (Al$_2$O$_3$, Cr$_2$O$_3$, TiO$_2$) or carbides (SiC, WC, Cr$_3$C$_2$), or diamond particles into metallic matrix improves significantly mechanical properties such as hardness and wear resistance. Ni-, NiCo- or Cu-based coatings incorporated with soft solid particles such as PTFE [220], MoS$_2$ [221], WS$_2$ [222], graphite [223], h-BN [224], single-wall carbon nanotube (SWCNT) [225], graphene nanoplatelet [226], Ti$_3$C$_2$ MXenes [227], SrSO$_4$ [159], and BaCr$_2$O$_4$ [150] can act as excellent self-lubricant layers, especially in precise mechanical parts and in slide bearings. The Cu-Sn, Cu-Sn-TiO$_2$, Cu-Sn-PTFE, and Cu-Sn-PTFE-TiO$_2$ coatings were electrodeposited in a pyrophosphate electrolyte with PTFE emulsion and TiO$_2$ sol, which led to a low friction coefficient of 0.1 and high wear resistance [228]. Sufficient solid lubricants embedded in the metallic-based coatings enable a significantly reduced friction coefficient by the formation of a self-lubricating tribofilm. As compared with plasma nitriding and sputtering deposition of thin, precious metals on superalloys, the electrodeposition process is a simple and economical method. Recently, with the emergence of compositionally- and hydrodynamically modulated layer coatings, the possibility of slow-release coatings for semi-continuous lubrication and modification of diffusion coatings by heat treatment have received considerable interest in composite coatings and surface functionality technologies [219].

As mentioned in Section 5.3, some oxometallates such as alkaline earth sulfates and chromates were developed as effective solid lubricants for sliding and rolling contact components. However, oxometallate-containing composites/coatings produced with the above-mentioned methods are generally associated with powder metallurgy processes such as spark plasma sintering and hot pressing, which may result in a possible chemical reaction between oxometallates and oxide ceramics or decomposition of oxometallates at elevated
temperatures. Thus, it is quite noteworthy to find new low-temperature routes, such as electrodeposition, to fabricate oxometallate-containing composite coatings, not only for cost reduction. Electrodeposition has been employed to produce Ni-SrSO$_4$ composite coatings on superalloy Inconel 718 from a Watts electrolyte containing a SrSO$_4$ suspension. Figure 16 shows the friction coefficients of the Inconel 718 substrate, pure nickel coating, and Ni-SrSO$_4$ composite coatings in sliding against a SAE52100 steel ball. As compared with the pure nickel coating and the substrate, the composite coating incorporated 6.83 vol.% SrSO$_4$ exhibits the enhanced microhardness of 450 HV$_{0.1}$, a low friction coefficient of 0.31, and a small wear rate of $5.3 \times 10^{-4}$ mm$^3$/Nm in sliding against bearing steel ball. The improvement in tribological properties of composite coatings results from the enhanced hardness and the resistance to plastic deformation and scuffing wear. Therefore, electrodeposited Ni-SrSO$_4$ composite coating is a potential self-lubricating material to improve the friction and wear properties of superalloys from room temperature to high operating temperatures for applications in aircraft gas engines, space mechanisms, petrochemical equipment, and offshore industries [159].

![Figure 16. Friction coefficients of Inconel 718 substrate, pure nickel coating and Ni-SrSO$_4$ composite coatings in sliding against SAE52100 steel ball. Reproduced with permission from Reference [159], Copyright © Elsevier B.V. 2009.](image)

In our previous work, electrodeposition was also employed to produce Ni-based chromate-containing composite coatings on Inconel 718 superalloy to improve friction and wear properties [150]. One challenge to depositing these coatings is that BaCrO$_4$ is soluble in an acid electrolyte solution, and a low concentration of Cr$^{6+}$ in the electrolyte significantly decreases the current efficiency of the deposition process, where only a little electrode current density can be assigned to the reduction of Ni$^{2+}$. In this case, the reduction reaction from Ni$^{2+}$ to metallic Ni at the cathode is very hard to push forward, and then the Ni-BaCrO$_4$ composite coating is difficult to achieve by electrodeposition. However, acid-insoluble BaCr$_2$O$_4$ particles instead of BaCrO$_4$ introduced into the nickel matrix are able to form a Ni-BaCr$_2$O$_4$ composite coating on Inconel 718 superalloy from a Watts electrolyte containing a BaCr$_2$O$_4$ suspension. Figure 17 shows the friction coefficients of the Inconel 718 substrate, pure nickel coating, and Ni-BaCr$_2$O$_4$ composite coatings incorporated with different volume fractions of BaCr$_2$O$_4$ in sliding against an Al$_2$O$_3$ ball. The Ni-16.6 vol.% BaCr$_2$O$_4$ composite coating exhibits a distinctly low friction coefficient of 0.31 and a small wear rate of $2.79 \times 10^{-6}$ mm$^3$/Nm as a comparative study with pure nickel coating and the substrate. The wear resistance of the electrodeposited Ni-BaCr$_2$O$_4$ composite coating was further improved by increasing the BaCr$_2$O$_4$ content. The improvements in tribological properties of Ni-BaCr$_2$O$_4$ composite coatings are attributed to both the
enhanced microhardness and self-lubricity of chromate-containing films formed on the sliding surface [150].

![Figure 17](image)

Figure 17. Friction coefficients of Inconel 718 substrate, pure nickel coating and Ni-BaCr$_2$O$_4$ composite coatings incorporated with different volume fractions of BaCr$_2$O$_4$ in sliding against Al$_2$O$_3$ ball. Reproduced with permission from Reference [150], Copyright © Elsevier B.V. 2011.

Electrodeposition was employed to produce the composites comprising MoS$_2$ on vertically aligned carbon nanotubes (CNTs) films, and a low friction coefficient of 0.03 and a wear rate of $10^{-13}$ mm$^3$/(Nm) were achieved even at 300 °C, which were about 2 orders of magnitude better than those of MoS$_2$-based nanoparticulate coatings [229].

6. Challenges Highlighted in High Temperature Solid Lubrication Applications

Industrial applications for high-temperature lubrication include not only the aerospace (sliding and rolling contact bearings, seals, and gears in the propulsion system of reusable launch vehicles, and precision satellite and spacecraft components), and aviation industries (bearing, abradable seals), but also other industries (hot metal forming process, high-speed dry machining process, side dams for the thin-strip steel casting process, pantograph contact strip for electric railways, cylinder wall/piston ring in diesel engines and automotives, as well as various furnace components) [2,8].

6.1. Bearings for Advanced Propulsion Systems

High-temperature bearings that must be self-lubricated include bearings of the shuttle, air foil bearings, rolling bearings of a gas turbine engine, rudder bearings of supersonic aircraft, bearings of adiabatic diesel engines, etc. The classic failure mode of rolling-bearing, which accommodates relative motion primarily by the action of rolling with a small slip (sliding) enabling it to carry high contact stresses, is fatigue spalling under cyclic contact stressing. However, when the rolling surface contains substantial sliding, severe adhesive wear, commonly called scuffing or smearing, worsens the surface damage. For bearings that operate at high temperatures (up to 1200 °C) and/or at high speeds with high precision, refractory materials and compounds such as alumina, yttria-stabilized zirconia, and silicon nitride are considered. Therefore, adaptive hard and soft multilayered coatings must be considered for long-lifetime rolling-element bearings at elevated temperatures.

One example is the lubrication application for advanced engines in hypersonic vehicles, which requires the lubricity and reliability of solid lubricants from ambient temperature to above 1000 °C during high-Mach-number (larger than 5) flights [3]. Air foil bearings in high Mach engines are designed to operate under heavy loads in the air at 900 °C or even higher.
The lubrication demands are challenging at different contact stages, such as start-up at room temperature, acceleration, or deceleration during operation, and shutdown at elevated temperatures, which may be well-suited for developing thermally stable oxide lubrication.

6.2. Seals Components for Advanced Propulsion Systems

Sliding of the seal surface occurs in a direction normal to the leakage flow under even no lubrication condition (e.g., in gas-path components such as a turbine or compressor blade tip). Seal components are subjected to adhesive wear, abrasive wear, corrosive wear, fatigue wear, and blistering during operation. Abradable (rub-tolerant) seals are used at the rotor-stator interface in the compressor and turbine sections of gas turbine engines to maintain the close tolerance without catastrophic failure. Plasma-sprayed Ni-Cr-bonded chrome carbide and tungsten carbide coatings are usually applied as abradable coatings in sliding against rub-tolerant plasma-sprayed chrome oxide coatings to compressor and turbine castings [19].

New technologies and approaches are urgently needed to explore high-temperature sealing materials and technologies for next-generation advanced propulsion systems, particularly in hypersonic vehicles, to prevent damage from both hot engine combustion gases and the friction heat of the atmosphere. In this case, sliding seal components will need to operate and resist environmental damage from an extreme temperature higher than 1000 °C, water vapor, oxidation, or reduction. Previous work on both alumina-boria-silica and complex carbide-oxide fiber materials demonstrated their potential use as high-temperature seal materials, although their sliding durability was still limited due to the reactive nature of boria and the oxidation/abrasion mechanism of carbide constituents, especially at elevated temperatures [12].

6.3. Hot Metal Forming Process

Components used in the hot metal forming industry such as molds, rolls, tools, and dies usually operate under conditions of extremely high temperature (strip/pipe/profile temperature up to 1100 °C for rolling, up to 1200 °C in hot extrusion) and mechanical (more than 1.0 GPa) dynamic loads and high velocity (10 m/s for rolling), which results in severe plastic deformation wear and fatigue [176]. Different processes, such as sheet forming, hydroforming, warm forming, superplastic forming, deep drawing, hot stamping, and forging, are used to produce aluminum parts for the automotive industry. However, in order to prevent direct metallic contact, seizure, and galling by lowering the friction stresses, all these processes require proper lubrication [230]. Reducing the friction coefficient is very important in a metal-forming process in order to prolong the lifetime of the tool, enhance the formability of the material, and improve the quality of the finished product.

Layered graphite has been widely used for the cold and hot forming of aluminum; however, dark stains remaining on the surface of the formed parts usually lead to additional grinding or polishing. $h$-BN has not only essentially the same lubricating mechanism as graphite but also high thermal stability, low thermal expansion, and good thermal shock resistance, although its lubricity is inferior to graphite and MoS$_2$. Therefore, when lubrication at elevated temperatures or cleanliness of working environments is of critical concern, a clean $h$-BN solid lubricant is capable of successfully replacing dirty graphite or MoS$_2$ to lubricate the surface of the formed parts during metalworking processes [60]. The stability of the lubrication film and the resulting surface quality depend mainly upon the particle size and concentration of $h$-BN powders [69]. Besides $h$-BN, a variety of high-temperature lubricants of inorganic additives such as borate-, silicate-, and polyphosphate salts exhibit suitable lubrication properties, which helps to reduce material loss and energy consumption and finally ensure the quality control and dimensional stability of the hot-formed components [176].
6.4. High-Speed Dry Machining

Machining technology is closely related to the development of high-temperature cutting tool materials with hard and wear-resistant surfaces such as nitride-, carbide-, or carbonitride-based monolithic and multilayered hard coatings. In consideration of environmental protection requirements and the elimination of costs for coolant media recirculation and utilization, cutting and milling operations gradually shift to high-speed dry machining without any oil-based coolant used. Surface functionality design in nanostructure and chemistry is urgently needed for a modern intelligent machining operation, which endows hard coatings with synergistic self-adaptation to lubricate up to 1000 °C and assure surface chemical and structural reversibility during thermo-mechanical cycling to maintain high cutting performance [30].

High-quality precision components can be rapidly produced by high-speed dry machining at the tip of the cutting edge. The material removal rate depends mainly upon allowable feeding rates, tool speeds, and cutting temperatures. Face or crater wear from chip motion and flank wear from the rubbing action are the main failure modes for a cutting tool, which is significantly influenced by a built-up edge generated from highly strained and hardened fragments. Therefore, the resistance to tool wear depends mainly upon the hot hardness, strength, fracture toughness, chemical stability, and reactivity of tool materials [19].

Cemented carbide or carbonitride tool materials are composed of granular graded transition metallic carbide or carbonitride particles of tungsten, titanium, tantalum, niobium, and other refractory metals bonded with cobalt, nickel, and molybdenum, and are produced by powder metallurgy processes. Various single or multilayered hard coatings of ceramic materials such as TiN, Al₂O₃, TiCₓNₓ, TiVOₓNᵧ, TiAlCrSiN, doped DLC, or diamond are deposited on cemented-carbide substrates to increase tool life by ion plating, magnetron sputtering, or chemical vapor deposition. The properties of these hard nitride coatings are generally modified with various elements, such as aluminum, silicon, chromium, and vanadium, to provide additional properties such as thermal stability, low adhesion, and even easy-to-shear [231]. Mo-N- and W-N-based coatings are designed for induced oxidation during dry machining operations and create wear-resistant and lubricious oxide phases such as molybdates and tungstates upon heating [232]. Multi-arc ion plating was employed to deposit VAICN-Ag nanocomposite and VAICN/VN-Ag multilayered coatings on Inconel 718 alloy, and the synergistic lubrication effect of both AgVO₃ and V₂O₅ Magneli phases was evaluated to verify their contributions to the lowest friction coefficient of about 0.18 for VAICN-Ag coating at 600 °C [233]. A self-lubricating WC-Co-Cu-BaF₂/CaF₂ wear-resistant coating with a friction coefficient of only 0.02 was fabricated for the application of machining tools [195]. CaF₂ is filled into the micro-texture of the rake face of the tool to form a self-lubricating film to improve the high-temperature machining performance, especially at elevated temperatures above 450 °C [234]. A self-lubricating Ti(C,N) cermet material modified with a multilayer core-shell microstructure exhibited enhanced wear-resistance and cutting performance [235]. In order to improve cutting performance, the benefits of Al(OH)₃ shell on CaF₂ solid lubricant were validated on hot-pressed Al₂O₃/Ti(C,N) ceramic tools by incorporating with ZrO₂ whiskers-reinforced CaF₂@Al(OH)₃ [236]. Nowadays, hard nanocomposite or multilayered nitride coatings with adaptive surface texturing lubrication will be developed as next-generation cutting tool coating materials.

6.5. Electric Contacts for Electric Railways and Transmission in Space

The coupling action between frictional and electric contact must be taken into account when sliding and rolling electrical contacts are involved, such as pantograph contact strips for electric railways and slip-roll rings for electric transmission in space. In machines that utilize a commutator, electrical brushes must be able to realize the commutation function when the load current is transferred to the external circuit. In the case of a pantograph contact strip, the selected materials must be able to operate at a rated current of 1000 A, a rated voltage of 25 kV, a sliding speed of 300 km/h and an applied load of 70 N with
an allowable wear loss of only 1 mm/10,000 km) [4]. Arc discharge attack and high-
temperature mechanical wear are the key factors influencing the high-power and long-
lifetime performance of current collectors. Electric brush wear is generally due to adhesion
and particle transfer, and in some circumstances, fatigue and fracture are caused by the
mechanical impact. Most electric brushes are comprised of graphite-based materials such as
electrographite, carbon-graphite, and metal-graphite due to their low friction and wear and
high electrical conductivity, which are fabricated either by conventional powder metallurgy
process or by infiltration of porous graphite with molten metals (e.g., Ag, Cu) and other
additives [19]. The metallized MAX phase or carbon contact strip is designed for electric
railways, and the noble metal-containing slip-roll ring is employed for electric transmission
in space mechanisms. Graphene-reinforced copper matrix composites exhibit an enhanced
tensile strength of 253 MPa, a low friction coefficient, and a small wear rate due to excellent
comprehensive properties such as self-lubricity, mechanical reliability, and chemical and
thermal stability of graphene. Examples of graphene-reinforced copper matrix composites
exhibit a service lifetime of electrical contact performance of 10 times longer than that of
commercially available pure copper and are almost comparable to the CuAg20 contacts
used in electrical engineering systems [237].

6.6. Refractory Side Dams for Thin-Strip Steel Casting Process

With the development of the thin-strip steel casting process, more requirements for
two fixed refractory side dams are put forward associated with extreme circumstances such as
mechanical stress (pressed against the water-cooled rollers), high-temperature corrosion,
and thermal shock (due to contact with liquid steel), and wear (due to contact with nickel-
coated copper rollers and steel strip) [71–73]. The sides of the rollers must be sealed to block
the overflow of molten steel by high-performance side dams [53,70]. BN-based materials
incorporated with \(m\)-ZrO\(_2\) and SiC are employed as the best side-dam materials for twin
roll strip steel casting and as a break ring for horizontal continuous casting, as contrasted
with other candidate materials such as sialon-BN, Si\(_3\)N\(_4\)-Al\(_2\)O\(_3\)-BN, TiN-MgO-BN, AIN-BN,
mullite-AIN-BN, Si\(_3\)N\(_4\)-AIN-BN, AlON-BN, and C\(_7\)-SiO\(_2\)-BN [53,73].

7. Conclusions

Understanding the friction and wear behavior of various solid lubricants and self-
lubricating composites under extreme environments is of significant importance for differ-
ent industrial applications, such as advanced propulsion systems in aerospace and aviation,
nuclear power engines, automotive, metal processing (cutting, forming, forging), metal-
lurgy, electric railways, etc. A survey has been made of various compounds which could be
considered environmentally compatible solid lubricant materials. The main groups of solid
lubricants are polymers, soft metals, laminar solids, chemically stable fluorides, binary or
ternary oxides, chromates, sulfates, and combinations of various solid lubricants.

1. The noble metals such as Ag and Au offer good lubricity due to enhanced ductility
and plastic deformation over a wide temperature range. The polymer composites
containing PTFE or polyimides provide lubrication with the lowest temperature
capacity, up from 300 to 350 °C.

2. MoS\(_2\)/WS\(_2\) are able to form a transfer film and generate excellent lubrication in
a vacuum and dry N\(_2\), while a graphite-like transfer film from graphite and DLC
provides lubrication in moist air. Layer-lattice solid lubricants such as graphite, MoS\(_2\),
and graphite fluoride generate structural degradation such as oxidation or dissociation
at certain temperatures, as well as the complex chalcogenides of Cs\(_2\)MoOS\(_3\), Cs\(_2\)WOS\(_3\),
and ZnMoOS\(_3\).

3. CaF\(_2\) and BaF\(_2\)/CaF\(_2\) eutectic are chemically stable non-layered inorganic compounds
under oxidizing environments, which exhibit low shear strength and easy film-
forming ability to provide good lubricity from 500 to 900 °C.
(4) Alkaline earth chromates of BaCrO$_4$ and BaCr$_2$O$_4$, and sulfates of BaSO$_4$ and SrSO$_4$ and their solid solutions show very good thermal stability and exceptional promise for lubricity over a wide temperature range.

(5) For extreme temperature circumstances, oxide lubrication is the focus of future studies. A new approach to solving low-temperature brittleness in oxide lubrication is to reduce their grain size to a few nanometers. In this case, plastic deformation in large part results from grain boundary sliding or grain rotating and only a minor contribution is associated with dislocation activity in ultrafine grains.

(6) Self-lubricating composites/coatings have been developed by a variety of material preparation techniques, which include powder metallurgy, physical/chemical vapor deposition, thermal spraying, electrodeposition, laser cladding, and additive manufacturing.

(7) Synergetic effects of different solid lubricants are widely explored for humidity-, temperature-, vacuum- or load-adaptive tribological applications. The underlying adaptive mechanisms are associated with environmental-assisted oxidation or interfacial tribo-reaction to form easy-to-shear and low-melting-point binary and ternary compounds, temperature-activated diffusion or melting of soft metals, and thermomechanically induced softening or surface self-glazing.

(8) The challenge associated with wide-range solid lubrication is the reversibility of the humidity-, temperature-, vacuum-, or load-adaptive tribological surfaces over multiple thermal cycles occurring in various engineering applications. Various approaches are postulated for adaptive multilayered coatings and surface multifunctional design, such as bionic composting, tunable surface texturing, and tribo-reaction of oriented lubricants. Temperature-adaptive composites/coatings exhibiting diffusion-, melting-, oxidation-, or tribo-reaction-limiting lubrication are developed over multiple thermal cycles through microlaminate architectures to activate the functionality of various solid lubricants.

(9) Adaptive solid lubrication design that can operate on earth and in space from room temperature to 1000 °C or even higher would be considered a breakthrough, which would increase air and space vehicle lifetime and performance. The challenges in high-temperature solid lubrication applications such as sliding and rolling contact bearings, seal components in advanced propulsion systems, hot-metal forming, high-speed dry machining, pantograph contact strips for electric railways, and side dams for thin-strip steel casting are highlighted. Microstructurally engineered combinations of solid lubricants will be of significant importance for the development of advanced lubrication systems under extreme environments of low/high temperature, high pressure, high chemical reactivity, and ultrahigh vacuum.

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