

Thermal-barrier coatings for more efficient gas-turbine engines

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Guest Editors

Gas-turbine engines used in transportation, energy, and defense sectors rely on high-temperature thermal-barrier coatings (TBCs) for improved efficiencies and power. The promise of still higher efficiencies and other benefits is driving TBCs research and development worldwide. An introduction to TBCs—complex, multi-layer evolving systems—is presented, where these fascinating systems touch on several known phenomena in materials science and engineering. Critical elements identified as being important to the development of future TBCs form the basis for the five articles in this issue of *MRS Bulletin*. These articles are introduced, together with a discussion of the major challenges to improved coating development and the rich opportunities for materials research they provide.

Introduction

Thermal-barrier coatings (TBCs) are refractory-oxide ceramic coatings applied to the surfaces of metallic parts in the hottest part of gas-turbine engines (**Figures 1 and 2**), enabling modern engines to operate at significantly higher gas temperatures than their predecessors (see recent reviews^{1–6}). Gas-turbine engines, used to propel aircraft and to generate electricity, are Carnot engines where their efficiency and core power are directly related to the gas temperature entering the turbine section.^{7,8} Further increases in the energy efficiency of gas-turbine engines, both to increase the electricity output and, for jet engines, the thrust-to-weight ratio and durability, will rely on further improvements in TBCs. At the same time, as gas temperatures are increased in the pursuit of higher engine efficiency, there are new challenges to existing TBCs.

To place this in context, gas-turbine engines are a \$42 billion industry worldwide (2010), with ~65% of the sales accounting for jet engines and the remainder land-based engines for electricity generation.⁹ The latter, fueled by natural gas or liquid fuels, produce ~25% of all electricity in the United States and ~20% worldwide (2010).¹⁰ With the anticipated worldwide growth of electricity demand and the recent discovery of vast shale gas resources, the number of gas-turbine engines in service will inevitably grow in the coming decades.⁹ Similarly, airline traffic is expected to double in the next 20 years,¹¹ while at the same time, there is a need to reduce high-altitude NO_x

pollution produced by jet engine exhausts.¹² Together, these developments will require continued innovation in gas turbine technology and high-temperature engine materials, including TBCs and associated technologies.

Many engineering design factors influence the overall efficiency of gas-turbine engines, but a major step in increasing engine temperature and engine efficiency was the introduction of TBCs. Typically made of ~7 wt% Y₂O₃-stabilized ZrO₂ (7YSZ) ceramics, TBCs provide thermal insulation to the metallic/superalloy engine parts. These parts include the combustor (**Figures 1 and 2**); stationary guide vanes, rotating blades (**Figure 1**), blade outer air-seals, and shrouds in the high-pressure section behind the combustor; and afterburners in the tail section of jet engines. As illustrated in **Figure 3**, the gas-temperature increase facilitated by the use of TBCs, in conjunction with innovative air-cooling approaches, has been much greater than that enabled by earlier materials development, including the development of single-crystal Ni-based superalloys.

Originally, TBCs were introduced to extend the useful life of stationary engine parts such as the combustor, but in the late 1980s, TBCs were first used on rotating blades.¹³ However, TBCs were not “prime reliant”; in other words, the ceramic coating was not considered in the design of the temperature capability of the underlying metal parts. Today, TBCs are critical components in gas-turbine engines, and because the gas temperatures are typically higher than the melting point of

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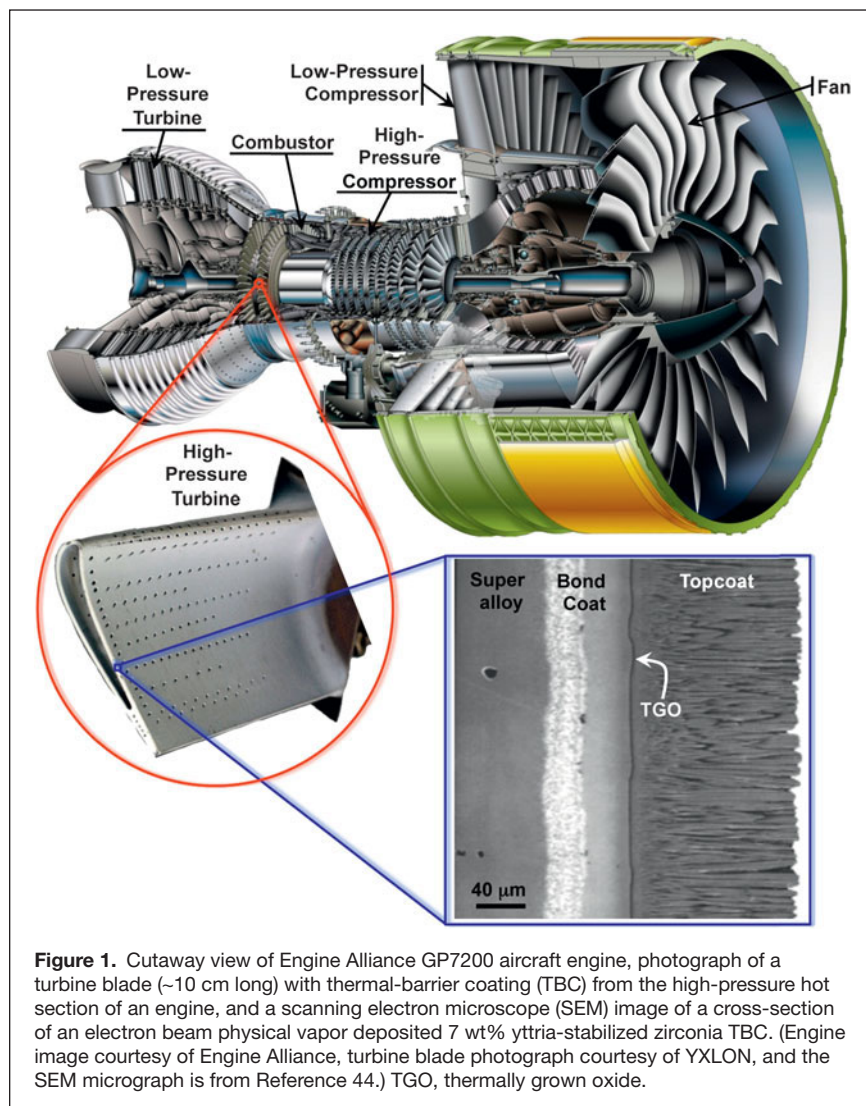


Figure 1. Cutaway view of Engine Alliance GP7200 aircraft engine, photograph of a turbine blade (~10 cm long) with thermal-barrier coating (TBC) from the high-pressure hot section of an engine, and a scanning electron microscope (SEM) image of a cross-section of an electron beam physical vapor deposited 7 wt% yttria-stabilized zirconia TBC. (Engine image courtesy of Engine Alliance, turbine blade photograph courtesy of YXLON, and the SEM micrograph is from Reference 44.) TGO, thermally grown oxide.

the underlying metal parts, any TBC failure can endanger the engine.¹⁴ Furthermore, because of the coupled diffusional and mechanical interactions between the oxide ceramic coating and the underlying alloys at these high temperatures, it is essential to consider TBCs as a complex, interrelated, and evolving material system, consisting not only of the oxide ceramic coating (topcoat) itself but also the underlying superalloy engine part, and two other layers in between. These include a metallic bond-coat layer that is more oxidation resistant than the superalloy, and a thin, thermally grown oxide (TGO) layer that forms between the topcoat and the bond coat as result of bond-coat oxidation in-service. The bond-coat composition is designed to result in a TGO made of α - Al_2O_3 —a mechanically robust, effective barrier to oxygen diffusion. **Figure 4** illustrates this multilayer structure in a typical TBC system.

During service, several kinetic processes occur in parallel. Interdiffusion between the bond coat and the underlying superalloy occurs, driven by chemical potential gradients; Al diffuses from the bond coat to form the TGO; and microstructural,

chemical, and phase changes occur in all the materials, including in the ceramic topcoat itself, changing their very properties. Since all of these are thermally activated processes, the rates at which they occur are expected to increase exponentially with temperature, albeit with different activation energies. Furthermore, the processes generally lead to degradation and failure of the coating.

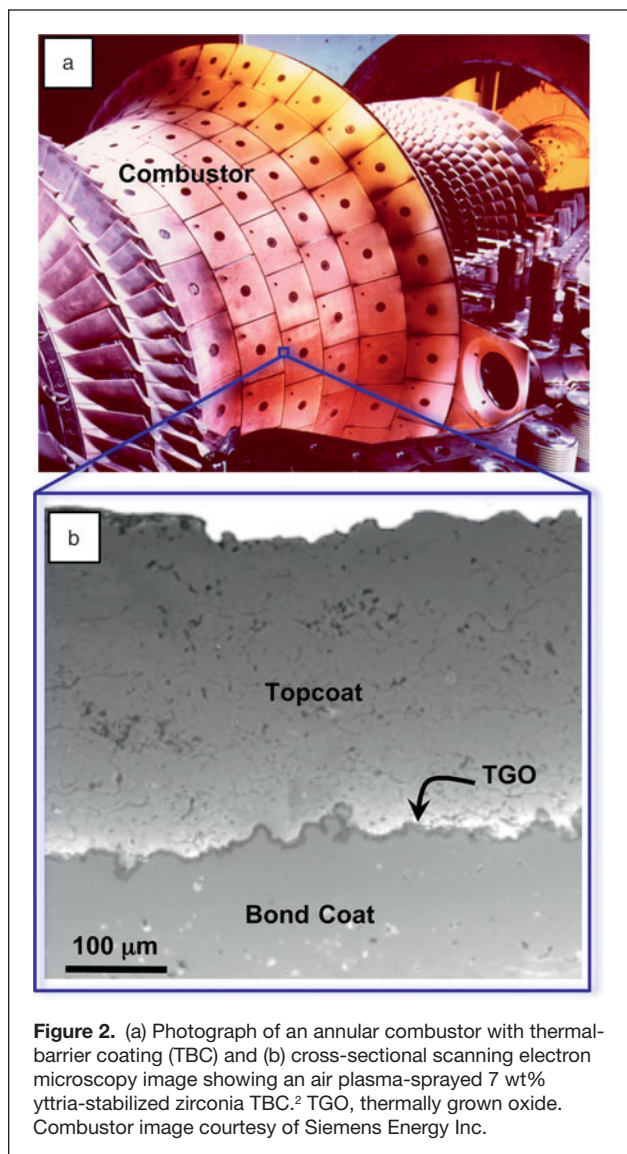
TBCs are also multifunctional: they must provide thermal insulation to protect the underlying superalloy engine parts, have strain compliance to minimize thermal-expansion-mismatch stresses with the superalloy parts on heating and cooling, and must also reflect much of the radiant heat from the hot gas, preventing it from reaching the metal alloy. Furthermore, TBCs must maintain thermal protection for prolonged service times and thermal cycles without failure. Typically, these times are 1000s of hours for jet engines being cycled numerous times between a maximum temperature of ~1300°C and room temperature (takeoff/landing and on-ground), and 10,000s of hours for power-generation engines with fewer thermal cycles (maintenance shut-downs). However, the latter are now being increasingly employed to stabilize the electric grid connected to renewable sources (wind, solar), and, thus, these engines experience more frequent cycles to compensate for the inherent intermittency of renewables. TBCs need to do this without separating from the engine parts while also withstanding extreme thermal gradients (~1°C μm^{-1}) and energy fluxes (~1 MW m^{-2}). Not only are these demands extremely exacting but also are often conflicting: TBCs must have both

low thermal conductivity and low weight; they must remain intact while withstanding large stress variations, both due to heating and cooling as well as under thermal shock; they must be chemically compatible with the underlying metal and the TGO; and they must operate in an oxidizing environment at maximum pressures of ~10 atmospheres and maximum gas velocities exceeding Mach 1.

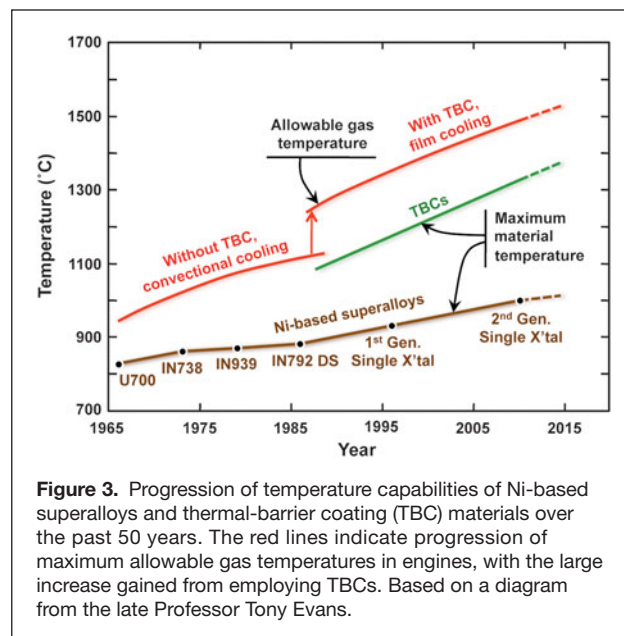
These demands and the desire to operate at higher temperatures reliably for longer times are driving new TBC innovations based on exploration of the underlying materials, processing sciences, and mechanistic understanding of degradation/failure and its mitigation. Several but not all of these key areas are highlighted in this issue of *MRS Bulletin*.

In this issue Ceramic topcoat processing

Sampath et al. describe the oxide ceramic topcoat deposition processes and microstructures. Unlike more traditional thin films used in microelectronics and in materials-growth studies,



these coatings are of intermediate thickness (100 μm to 1 mm), and they must be deposited at a high rate to incorporate porosity. Typical TBC porosity is $\sim 15\%$, which is essential for high strain compliance and reduced thermal conductivity, yet the TBCs need to be mechanically robust to resist fracture, erosion, and foreign object damage (FOD). Furthermore, TBCs need to be deposited on complex-shaped parts with highly curved surfaces, and at the same time the TBCs must have reproducible thermal and mechanical properties. Currently, TBCs are deposited by air plasma-spraying (APS)¹⁵ or by electron beam physical vapor deposition (EBPVD).¹⁶ Typically, the low-cost APS method is used to deposit TBCs on stationary engine parts (combustor, shroud, vanes), whereas EBPVD TBCs are used on the most demanding hot-section parts in jet engines such as blades and vanes. Today, both stationary and rotating hot-section parts in electricity-generation engines, which tend to be much larger than those in jet engines, use APS TBCs. The microstructures of APS and EBPVD TBCs are vastly different (see Figures 1 and 2),

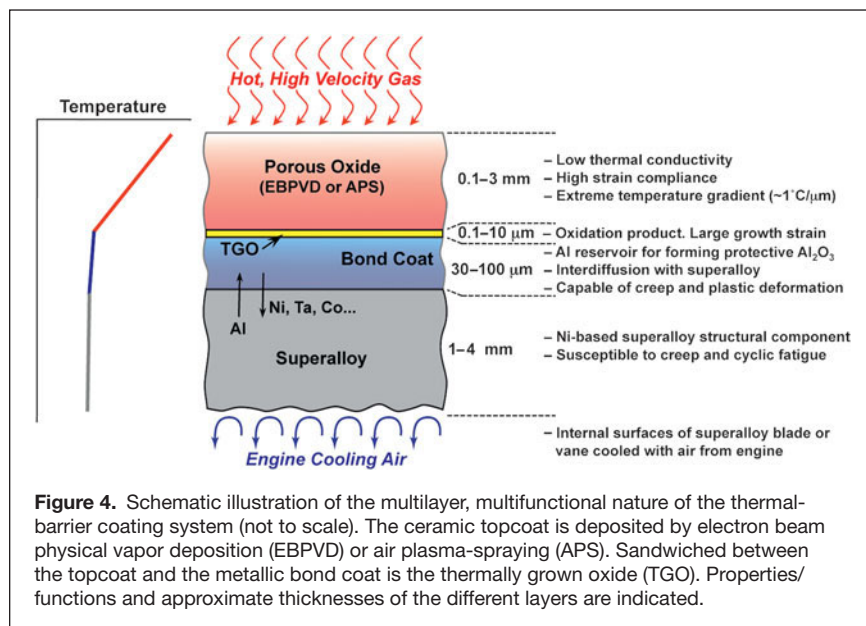


and each offer different advantages in terms of properties and performance. While processing science and technology of depositing 7YSZ TBCs is well established, the widespread introduction of improved TBCs of alternative compositions will ultimately depend on their reproducible and successful deposition commercially. Alternative TBC deposition methods with added advantages and versatilities are also being pursued.^{17–20}

TBCs testing and evaluation

An equally big hurdle to developing improved TBCs is the sheer complexity and variety of failure modes and their dependence on engine operating conditions. Superficially, the failures are similar; the coating spalls off the engine part exposing the underlying metal to rapid oxidation or melting. However, a variety of mechanisms can be responsible for the observed failure.¹ In some instances, the TGO grows to exceed a critical thickness and spalls off, causing TBC failure.^{21,22} In others, the initially flat bond coat and TGO undergo a complex morphological instability—“rumpling” (**Figure 5**)—causing local separations that grow in size with thermal cycling until linking up to form a spall.^{23–25} In still others, cavitation occurs in the bond coat and grows under thermal cycling.²⁶ In each of these cases, the failure location depends on the actual thermal and mechanical loading conditions in the coating. Typically, at moderate heat fluxes but high temperatures at the bond coat, failure is dominated by processes related to TGO formation and bond-coat inelastic behavior, where fracture occurs in the bond-coat/topcoat interface region. With increasing topcoat surface temperature and thermal-gradient, thermal-expansion mismatch starts to play a more dominant role, and fracture location typically shifts to within the ceramic topcoat.

In some cases, spallation occurs as a result of impact from particles (FOD) carried along with the hot, combusted gas.²⁷ In other cases, thermal-shock spallation of the coating can occur



during rapid cooling. In still other cases, fine sand and particulates ingested into the engine (including volcanic ash) can melt into a silicate glass at high temperatures ($>1200^\circ\text{C}$) on the TBC surface and wick into the TBC, decreasing its strain compliance and causing spallation upon cooling.

Both the crack driving force and the fracture resistance offered by the material also depend on the mechanical and thermophysical properties of the TBC system. These can change significantly with time at temperature. Also, these properties can be influenced significantly by the coating microstructure, which in turn is influenced by the process parameters used during the coating deposition. Even small variations of these parameters can lead to significant changes in coating microstructure, and thus to a wide variation in mechanical and thermophysical properties, resulting in a large scatter in system reliability, durability, and predictability. Thus, deterministic approaches for predicting TBC life can be often misleading, requiring the use of probabilistic approaches for predicting TBC life. From a manufacturing, as well as materials selection perspective, clarifying mechanisms of failure and being able to predict life remains an essential and central task in optimizing the coating system.

This has motivated the development of TBC testing methods and non-destructive evaluation techniques, especially under realistic conditions pertinent to engine operation, which can include high pressures (up to 10 atm.), high temperatures (up to 1400°C), steep thermal gradients (temperature differences up to 300°C), high gas velocities, and the presence of detrimental environmental species (e.g., water vapor, sand, ash, salt). The article by Vaßen et al. describes some of the common TBC failure modes and innovations in TBC testing and evaluation.

Topcoat ceramics

The majority of TBCs in use today are ZrO_2 -based having a composition containing ~ 7 wt% Y_2O_3 (7YSZ). Originally,

this ceramic was selected empirically based on its low thermal conductivity, high melting point, resistance to sintering, a demonstrated manufacturing capability for depositing it with constant composition, and long life in the resulting TBCs.^{13,28–30} Unlike the cubic ZrO_2 used in oxide fuel cells, oxygen sensors, and fake diamonds, which have higher Y_2O_3 content, 7YSZ is a metastable tetragonal phase (t').⁴ 7YSZ has been shown to have unusually high fracture toughness due to ferroelastic toughening.^{31,32} Unlike other transformation-toughened ZrO_2 -based ceramics, so-called “ceramic steels,”³³ used in bearings, cutting tools, and knives, the toughness in 7YSZ does not arise from the martensitic transformation (an irreversible and diffusionless collective movement of atoms) from the tetragonal to monoclinic phase but rather from reversible ferroelastic domain switching from one tetragonal variant to another when stressed.^{31,32} Also, unlike transformation

toughening, ferroelastic toughening can operate at high temperatures, typical of those at engine temperatures. High fracture toughness in TBCs is important not only for resisting impact and erosion but also spallation.

Despite these intrinsic advantages, there is a worldwide search under way for oxides with superior, high-temperature properties that could replace 7YSZ. Much of this activity is presently directed to identifying oxides with lower thermal conductivity,³⁴ as discussed in the article by Pan et al. Although the underlying physics of thermal conductivity in solids was firmly established more than 30 years ago, the challenge is to translate the concepts to identify prospective low conductivity compounds in terms of crystal structure and bonding, especially when little or nothing is known about the phonon (lattice wave) properties of almost all poly-ionic oxides. The bulk of heat transport in these oxides occurs via phonons, and their scattering governs the oxide thermal resistance. Fortunately, at high temperatures, the majority of lattice phonons can be expected to be fully thermally activated, so classical descriptions of thermal conductivity can guide the search for low conductivity oxides. This search has, for instance, revealed that natural superlattice structures have exceptionally low thermal conductivity^{35–37} as do oxides with a large number of ions per unit cell that also can exhibit extensive solid solution.³⁸ Another insight, gained from molecular dynamics simulations, is that in YSZ, phonons are highly delocalized and transport diffusively, akin to a phonon glass, despite the crystal perfection measured by x-ray diffraction.³⁹

Bond-coat alloys and oxidation

In many respects, the most stringent constraints are imposed on the bond coat. Its primary function is to provide a reservoir from which Al can diffuse to form a protective $\alpha\text{-Al}_2\text{O}_3$ TGO while maintaining cohesion with the TBC without reacting with it. Mechanics modeling⁴⁰ indicates that, ideally, the TGO

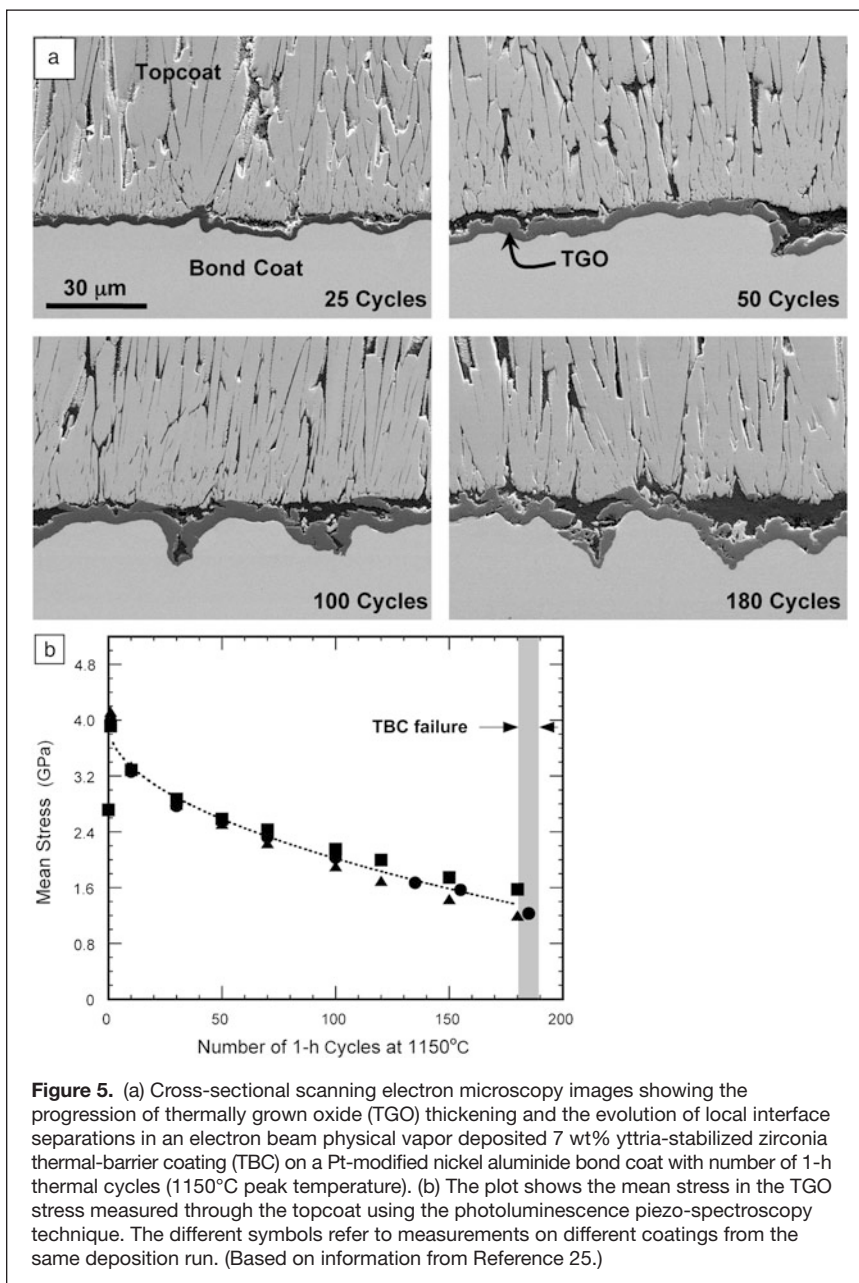


Figure 5. (a) Cross-sectional scanning electron microscopy images showing the progression of thermally grown oxide (TGO) thickening and the evolution of local interface separations in an electron beam physical vapor deposited 7 wt% yttria-stabilized zirconia thermal-barrier coating (TBC) on a Pt-modified nickel aluminide bond coat with number of 1-h thermal cycles (1150°C peak temperature). (b) The plot shows the mean stress in the TGO stress measured through the topcoat using the photoluminescence piezo-spectroscopy technique. The different symbols refer to measurements on different coatings from the same deposition run. (Based on information from Reference 25.)

should remain elastic to the highest temperatures and not creep to prevent “rumpling”^{23,24} or cavitation on thermal cycling²⁶ that can, in turn, lead to the development of local separations at the TBC interface (Figure 5).²⁵ At the same time, it has to operate at the highest temperature possible to minimize the amount of air used to cool the vanes and blades, without reacting with the underlying superalloy and melting. This presently implies that the maximum bond-coat temperature cannot be allowed to exceed ~1150°C. Currently, there are two main bond-coat alloys in use, a Ni-rich nickel aluminide and a compositionally more complex MCrAlY (M=Ni, Co+Ni, or Fe) alloy. While these are very different alloys metallurgically, the challenges are similar, as described in the article by Pollock et al.: how to minimize deformation at intermediate and operating temperatures, how

to minimize interdiffusion with the underlying superalloy to prevent the formation of brittle intermetallics, and how to deliver critical elements in addition to Al, such as Hf and Y, to the growing TGO to minimize its inelastic plastic deformation under thermal cycling.

Critical to understanding the performance of the TBC system is the formation, growth, and properties of the TGO that forms underneath the 7YSZ topcoat by oxidation of the bond-coat alloy (TBC microstructures are highly defective with porosity and cracks, and 7YSZ is an oxygen conductor, hence oxidation of the bond coat cannot be prevented). The bond-coat compositions are selected to form an α -Al₂O₃ TGO because it is the slowest growing oxide at high temperatures and forms an impervious, adherent layer with excellent mechanical integrity. This is important because TBC failure can occur when the TGO growth exceeds a critical thickness. The essential mechanics of this form of failure are similar to the origin of a critical thickness for the loss of coherence of epitaxial thin films, namely when the release of stored elastic strain energy in the growing film exceeds the fracture resistance.⁴¹ There are two contributions to the stress in the TGO, one is associated with the growth strain as new oxide is created at the grain boundaries of the TGO, and the other is the mismatch stress with the superalloy generated by differences in thermal expansion on cooling. The growth strain consists of two components: one that leads to a simple thickening and the other that motivates lateral expansion of the TGO that, in turn, drives out-of-plane instabilities as well as other mechanical responses.⁴⁰ The origin of the lateral growth strain is poorly understood but is generally attributed to the counter-diffusion of inward diffusing O²⁻ and outward diffusing Al³⁺, resulting in the plating out of new Al₂O₃ in the TGO grain bounda-

ries.⁴² There have been a limited number of measurements of the growth strain in the TGO absent the TBC itself using x-ray synchrotron sources⁴³ but not nearly enough to follow the evolution during oxidation or thermal cycling. More revealing have been non-contact measurements by photoluminescence piezo-spectroscopy of the strains measured through the topcoat.⁴⁴ In this technique, a laser beam is used to penetrate through the topcoat and excite the *R*-line luminescence from trace Cr³⁺ ions invariably present in the TGO. The local mean stress in the TGO is proportional to the frequency shift of the *R*-lines. This has enabled correlations to be mapped between luminescence shifts and the development of local damage as the bond coat and TGO rumple, as shown in **Figures 5** and **6**.^{23,25,45} There remain several important unresolved questions about the lateral

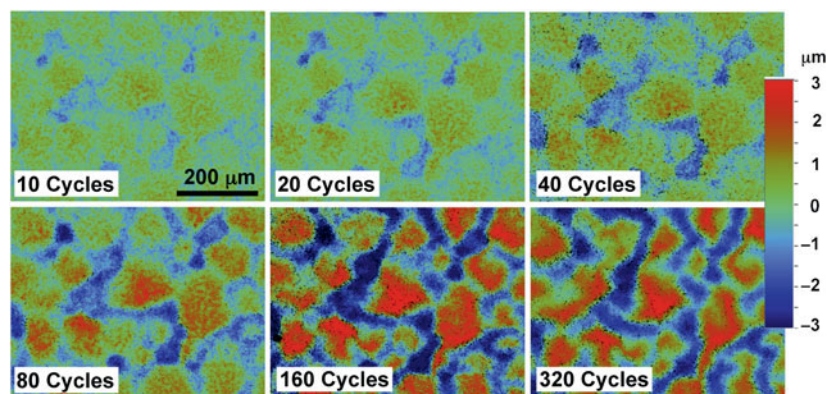


Figure 6. Topographic profilometer (optical) images (top view) of the identical area of an aluminized bond coat (without a ceramic topcoat) after polishing flat and then thermal cycled (1-h cycles) between room temperature and 1150°C for the cycles indicated. As is evident from the sequence of images, the magnitude of the rumpling surface instability increases with cycling, but the microstructural scale does not. The color scale at the right indicates the rumpling height variation. (Based on information from Reference 45.)

growth strain whose resolution could impact oxidation of other metallic alloys. These include how minor elements, at the ppm level and above, affect the growth and mechanical behavior of the TGO. Of particular interest are the elements Y, Zr, and Hf that segregate, on account of their large ionic radii, to the grain boundaries of the TGO. Among the key questions being raised are whether these elements alter the counter-diffusion along the TGO grain boundaries that creates the lateral growth strain and how they affect the high-temperature creep and plasticity of the TGO. It is known that rare-earth ions dramatically increase the creep resistance of alumina ceramics.⁴⁶

Attack by molten deposits and its mitigation

Higher engine temperatures are also creating new materials issues in ceramic topcoats, namely degradation of 7YSZ TBCs due to molten silicate deposits,^{47–52} formed by the ingestion of fine particulates from the environment (sand,⁴⁹ volcanic ash^{53,54}) (see the Levi et al. article in this issue). Because of the major components in the silicate glass formed, this phenomenon is commonly referred to as CMAS (calcium-magnesium-alumino-silicate) attack. This primarily affects high-performance jet engines on account of their higher maximum temperatures and electricity-generation engines in some locations, but it will likely affect more engines as operation temperatures are increased in pursuit of greater engine efficiencies. In the case of land-based electricity-generation engines, it is not always practical to filter out the finest particles that can be carried along with the input air and from alternative fuels such as syngas.^{55,56} It appears that wetting of TBCs by the molten CMAS glass, and dissolution/reprecipitation

of YSZ grains in that glass, contribute to the CMAS attack of 7YSZ TBCs.^{50,51} This manifests itself as continued penetration of the CMAS glass into the TBC and affects both APS and EBPVD TBCs alike. For example, **Figure 7** shows complete penetration of an EBPVD 7YSZ TBC by molten CMAS in a laboratory test.⁵⁷ Therefore, being able to mitigate CMAS attacks becomes an additional critical requirement for future TBCs.

While this area of TBC research is relatively new, important insights into CMAS attack and mitigation mechanisms have started to emerge. For instance, for a TBC ceramic to be highly effective against CMAS attack, wetting should be prevented, and/or it must readily interact with the CMAS to form a crystalline sealing layer arresting further penetration of the molten CMAS.^{51,58–63} By corollary, TBCs made from ceramics that are inert or stable in contact with molten CMAS may not be effective in resisting CMAS wetting and attack. Thus, tailoring of TBC compositions and microstructures for resistance against molten silicate deposits is becoming an important area of TBC research. An example of APS TBC of designed composition (7YSZ containing Al_2O_3 and TiO_2) resisting attack by molten ash from the Eyjafjallajökull volcano in a laboratory test is shown in **Figure 8a–b**.⁵⁴

Outlook

The sheer complexity of the interactions between the four principal layers and materials in the thermal-barrier coating (TBC) system—ceramic topcoat, thermally grown oxide, metallic bond coat, base superalloy—and their evolution with time at

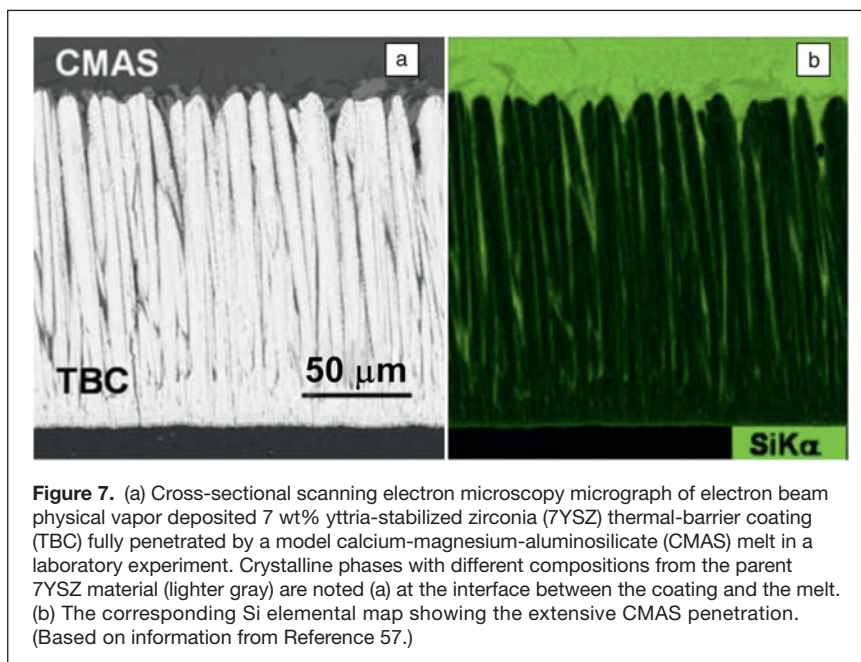


Figure 7. (a) Cross-sectional scanning electron microscopy micrograph of electron beam physical vapor deposited 7 wt% yttria-stabilized zirconia (7YSZ) thermal-barrier coating (TBC) fully penetrated by a model calcium-magnesium-aluminosilicate (CMAS) melt in a laboratory experiment. Crystalline phases with different compositions from the parent 7YSZ material (lighter gray) are noted (a) at the interface between the coating and the melt. (b) The corresponding Si elemental map showing the extensive CMAS penetration. (Based on information from Reference 57.)

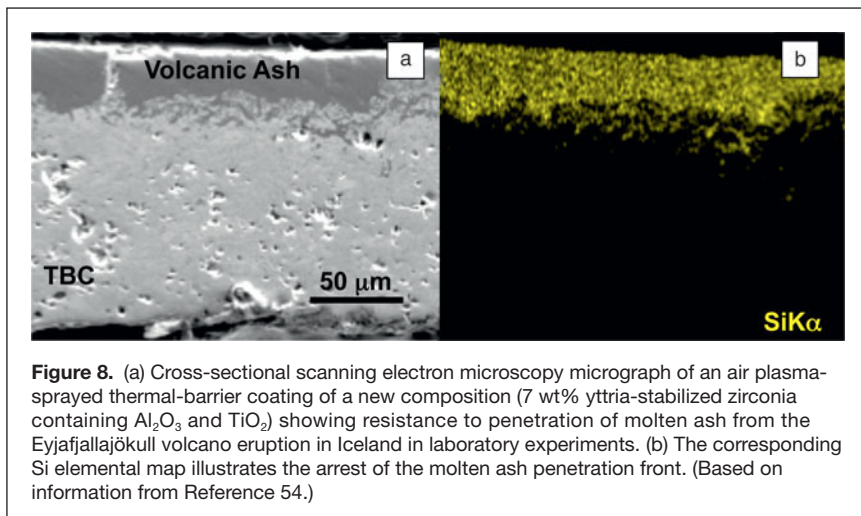


Figure 8. (a) Cross-sectional scanning electron microscopy micrograph of an air plasma-sprayed thermal-barrier coating of a new composition (7 wt% yttria-stabilized zirconia containing Al_2O_3 and TiO_2) showing resistance to penetration of molten ash from the Eyjafjallajökull volcano eruption in Iceland in laboratory experiments. (b) The corresponding Si elemental map illustrates the arrest of the molten ash penetration front. (Based on information from Reference 54.)

temperature make it essential that synergistic progress be made in all areas to enable TBCs to operate reliably at still higher temperatures in the future. Three major challenges stand out. The first is to increase the reproducibility of the coating deposition so that full temperature capabilities of existing TBCs can be utilized with greater confidence. Indeed, at present, engine designers only take into account about half of the possible temperature increase afforded by the thermal properties of current TBCs because of the lack of processing reproducibility. The second challenge is to have more comprehensive modeling of the evolution of the coating system and its failure, as well as a better description of the material properties, especially at high temperatures. This will require a concerted modeling effort spanning multiple length and temporal scales, supported by experimental validation that will greatly benefit the development and implementation of future TBCs. The third major challenge is tackling new issues that arise with higher temperatures. One such issue is radiative heat transport through the TBC. New approaches will be needed to reflect and/or scatter radiation and prevent it from reaching the metallic parts. Another important issue is calcium-magnesium-alumino-silicate (CMAS) attack and its mitigation. This will require new TBC compositions and microstructures that not only resist CMAS penetration but also meet a suite of other requirements at those higher temperatures.

While the use of TBCs has already resulted in dramatic improvements in the efficiency and the power output of gas-turbine engines, the challenges described in this and the companion articles present new materials research opportunities essential for turbine engine designers to take full advantage of recent advances in materials, processing, and reliability. This will become even more crucial in bridging the growing energy and transportation demands of society until large scale energy generation from renewable sources (solar, wind) becomes economically more viable. Indeed, because gas turbines already play a major role in electricity generation and aircraft propulsion, even minor improvements in engine efficiency will

have an immediate and significant positive impact on the overall energy portfolio of the world.

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References

1. A.G. Evans, D.R. Mumm, J.W. Hutchinson, G.H. Meier, F.S. Pettit, *Prog. Mater. Sci.* **46**, 505 (2001).
2. N.P. Padture, M. Gell, E.H. Jordan, *Science* **296**, 280 (2002).
3. C.G. Levi, *Curr. Opin. Solid State Mater. Sci.* **8**, 77 (2002).
4. D.R. Clarke, C.G. Levi, *Annu. Rev. Mater. Sci.* **33**, 383 (2003).
5. A.G. Evans, D.R. Clarke, C.G. Levi, *J. Eur. Ceram. Soc.* **28**, 1405 (2008).
6. R. Vaßen, M. Ophelia-Jarligo, T. Steinke, D. Emil-Mack, D. Stöver, *Surf. Coat. Technol.* **205**, 938 (2010).
7. W.W. Bathie, *Fundamentals of Gas Turbines—Second Edition* (Wiley, New York, 1996).
8. J.H. Perepezko, *Science* **326**, 1068 (2009).
9. L.S. Langston, *Mech. Eng.* **133**, 30 (2011).
10. US Energy Information Administration, *Annual Energy Review*; www.eia.gov/totalenergy/data/annual/index.cfm (accessed June 2012).
11. Federal Aviation Administration, *FAA Forecast Predicts Air Travel to Double in Two Decades*; www.faa.gov/news/press_releases/news_story.cfm?newsId=12439 (accessed June 2012).
12. International Civil Aviation Organization, ICAO Environment Report, www.icao.int/environmental-protection/Pages/EnvReport10.aspx (accessed June 2010).
13. U. Schulz, K. Fritscher, C. Leyens, M. Peters, W.A. Kaysser, *JOM* **49** (1997).
14. Anon., *The Economist* **290**, 60 (2009).
15. H. Herman, S. Sampath, R. McCune, *MRS Bull.* **25**, 17 (2000).
16. U. Schulz, C. Leyens, K. Fritscher, M. Peters, B. Saruhan-Brings, O. Lavigne, J.M. Dorvaux, M. Poulain, R. Mevrel, M.L. Caliez, *Aerosp. Sci. Technol.* **7**, 73 (2003).
17. N.P. Padture, K.W. Schlichting, T. Bhatia, A. Ozturk, B. Cetegen, E.H. Jordan, M. Gell, S. Jiang, T.D. Xiao, P.R. Strutt, E. Garcia, P. Miranzo, M.I. Osendi, *Acta Mater.* **49**, 2251 (2001).
18. P. Fauchais, G. Montavon, *J. Therm. Spray Technol.* **19**, 226 (2010).
19. A. Hospach, G. Mauer, R. Vaßen, D. Stöver, *J. Therm. Spray Technol.* **20**, 116 (2011).
20. K. von Niessen, M. Gindrat, *J. Therm. Spray Technol.* **20**, 736 (2011).
21. K. Chan, S. Cheruvu, R. Vishwanathan, *ASME Turbo Expo 2003, Atlanta* (2003), p. 591.
22. K.W. Schlichting, N.P. Padture, E.H. Jordan, M. Gell, *Mater. Sci. Eng., A* **342**, 120 (2003).
23. V.K. Tolpygo, D.R. Clarke, *Acta Mater.* **52**, 5115 (2004).
24. M. Wen, E.H. Jordan, M. Gell, *Surf. Coat. Technol.* **201**, 3289 (2006).
25. B. Heeg, V.K. Tolpygo, D.R. Clarke, *J. Am. Ceram. Soc.* **94**, S112 (2011).
26. V.K. Tolpygo, *Surf. Coat. Technol.* **2002**, 617 (2007).
27. X. Chen, R. Wang, N. Yao, A.G. Evans, K.J.J.W. Hutchinson, R.W. Bruce, *Mater. Sci. Eng. A* **352**, 221 (2003).
28. S. Stecura, *Am. Ceram. Soc. Bull.* **56**, 1082 (1977).
29. S. Stecura, H. Curt, US Patent No. 4,005,705 (1977).
30. R.L. Jones, in *Metallurgical and Ceramic Coatings*, K.H. Stern, Ed. (Chapman & Hall, London, UK, 1996).
31. A.V. Virkar, *Key Eng. Mater.* **153–154**, 183 (1998).
32. C. Mercer, J.R. Williams, D.R. Clarke, A.G. Evans, *Proc. R. Soc. London, Ser. A* **463**, 1393 (2007).
33. D.J. Green, R.H.J. Hannink, M.V. Swain, *Transformation Toughening of Ceramics* (CRC Press, Boca Raton, FL, 1989).

34. M.R. Winter, D.R. Clarke, *J. Am. Ceram. Soc.* **90**, 533 (2007).
 35. Y. Shen, D.R. Clarke, P.A. Fuierer, *Appl. Phys. Lett.* **93**, 102907 (2008).
 36. A. Chernatynskiy, R.W. Grimes, M.A. Zurbuchen, D.R. Clarke, *Appl. Phys. Lett.* **95**, 161906 (2009).
 37. T.D. Sparks, P.A. Fuierer, D.R. Clarke, *J. Am. Ceram. Soc.* **93**, 1136 (2010).
 38. Z. Qu, T.D. Sparks, W. Pan, D.R. Clarke, *Acta Mater.* **59**, 3841 (2011).
 39. P.K. Schelling, S.R. Phillpot, *J. Am. Ceram. Soc.* **84**, 2997 (2001).
 40. D.S. Balint, J.W. Hutchinson, *J. Mech. Phys. Solids* **53**, 949 (2005).
 41. L.B. Freund, S. Suresh, *Thin Film Materials: Stress, Defect Formation and Surface Evolution* (Cambridge University Press, Cambridge, UK, 2003).
 42. F.N. Rhines, J.S. Wolf, *Metall. Mater. Trans. A* **1**, 1701 (1970).
 43. P.Y. Hou, A.P. Paulikas, B.W. Beal, *JOM* **61**, 51 (2009).
 44. R.J. Christensen, D.M. Lipkin, D.R. Clarke, K. Murphy, *Appl. Phys. Lett.* **69**, 3754 (1996).
 45. S. Dryepondt, J. Porter, D.R. Clarke, *Acta Mater.* **57**, 1717 (2009).
 46. J. Cho, C.M. Wang, H.M. Chan, J.M. Rickman, M.P. Harmer, *Acta Mater.* **47**, 4197 (1999).
 47. J.L. Smialek, F.A. Archer, R.G. Garlick, *JOM* **46**, 39 (1994).
 48. F.H. Stott, D.J. DeWet, R. Taylor, *MRS Bull.* **19**, 46 (1994).
 49. M.P. Borom, C.A. Johnson, L.A. Peluso, *Surf. Coat. Technol.* **86-87**, 116 (1996).
 50. S. Kramer, J. Yang, C.G. Levi, C.A. Johnson, *J. Am. Ceram. Soc.* **89**, 3167 (2006).
 51. A. Aygun, A.L. Vasiliev, N.P. Padture, X. Ma, *Acta Mater.* **55**, 6734 (2007).
 52. A.G. Evans, J.W. Hutchinson, *Surf. Coat. Technol.* **201**, 7905 (2007).
 53. J. Kim, M.G. Dunn, A.J. Baran, D.P. Wade, E.L. Tremba, *ASME J. Eng. Gas Turbines Power* **115**, 641 (1993).
 54. J.M. Drexler, A.D. Gledhill, K. Shinoda, A.L. Vasiliev, K.M. Reddy, S. Sampath, N.P. Padture, *Adv. Mater.* **23**, 2419 (2011).
 55. J.P. Bons, J. Crosby, J.E. Wammack, B.I. Bentley, T.H. Fletcher, *ASME J. Eng. Gas Turbines Power* **129**, 135 (2007).
 56. A.D. Gledhill, K.M. Reddy, J.M. Drexler, K. Shinoda, S. Sampath, N.P. Padture, *Mater. Sci. Eng. A* **58**, 7214 (2011).
 57. M.-P. Bacos, J.-M. Dorvaux, S. Landais, O. Lavigne, R. Mevrel, M. Poulain, C. Rio, M.-H. Vidal-Setif, in *Aerospace Lab: The ONERA Journal*. ONERA, Chatillon, France, **3** (2011).
 58. M. Freling, M.J. Maloney, D.A. Litton, K.W. Schlichting, J.G. Smeggil, D.B. Snow, US Patent No. 7,455,913 (2008).
 59. S. Kramer, S. Faulhaber, M. Chambers, D.R. Clarke, C.G. Levi, J.W. Hutchinson, A.G. Evans, *Mater. Sci. Eng. A* **490**, 26 (2008).
 60. D.A. Litton, K.W. Schlichting, M. Freling, J.G. Smeggil, D.B. Snow, M.J. Maloney, US Patent No. 7,662,489 (2010).
 61. J.M. Drexler, K. Shinoda, A.L. Ortiz, D. Li, A.L. Vasiliev, A.D. Gledhill, S. Sampath, N.P. Padture, *Acta Mater.* **58**, 6835 (2010).
 62. J.M. Drexler, C.-H. Chen, A.D. Gledhill, K. Shinoda, S. Sampath, N.P. Padture, *Surf. Coat. Technol.* **206**, 3911 (2012).
 63. J.M. Drexler, A.L. Ortiz, N.P. Padture, *Acta Mater.* **60**, 5437 (2012). □



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