Review of the anatase to rutile phase transformation

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Abstract  Titanium dioxide, TiO₂, is an important photocatalytic material that exists as two main polymorphs, anatase and rutile. The presence of either or both of these phases impacts on the photocatalytic performance of the material. The present work reviews the anatase to rutile phase transformation. The synthesis and properties of anatase and rutile are examined, followed by a discussion of the thermodynamics of the phase transformation and the factors affecting its observation. A comprehensive analysis of the reported effects of dopants on the anatase to rutile phase transformation and the mechanisms by which these effects are brought about is presented in this review, yielding a plot of the cationic radius versus the valence characterised by a distinct boundary between inhibitors and promoters of the phase transformation. Further, the likely effects of dopant elements, including those for which experimental data are unavailable, on the phase transformation are deduced and presented on the basis of this analysis.

Background

Titanium dioxide, also known as titania, is of growing interest due to its proven ability to function as a photocatalyst and facilitate important environmentally beneficial reactions, such as water splitting to generate hydrogen and treatment of polluted air and water.

Titanium dioxide occurs as two important polymorphs, the stable rutile and metastable anatase. These polymorphs exhibit different properties and consequently different photocatalytic performances. Anatase transforms irreversibly to rutile at elevated temperatures. This transformation does not have a unique temperature and the processes that are involved in the transformation as well as the methods to inhibit or promote this transformation have not been reviewed comprehensively to date.

The present work aims to clarify the differences between the two main polymorphs of titanium dioxide, the nature of the anatase to rutile transformation, and the principles of controlling phase composition through the inhibition or promotion of the transformation of anatase to rutile.

Titania polymorphs

Titanium dioxide, the only naturally occurring oxide of titanium at atmospheric pressure, exhibits three polymorphs: rutile, anatase, and brookite [1–7]. While rutile is the stable phase, both anatase and brookite are metastable; the latter is difficult to synthesise and so is seldom studied [8]. Another five high-pressure phases of TiO₂ have been reported:

- TiO₂ II or srilankite, an orthorhombic polymorph of the lead oxide structure
- Cubic fluorite-type polymorph
- Pyrite-type polymorph
- Monoclinic baddeleyite-type polymorph
- Cotunnite-type polymorph

The stability of these phases has been discussed in several publications [4, 7, 9–12]. However, these are of minor significance for research and development applications.
Titania properties

Table 1 outlines the basic properties of rutile and anatase.

Titania applications

The primary application of titanium dioxide is as a white pigment in paints, food colouring, cosmetics, toothpastes, polymers, and other instances in which white colouration is desired [13]. The reason for this is the high refractive indices of rutile and anatase, which result in high reflectivity from the surfaces. Consequently, titanias of small particle size and correspondingly high surface areas are used owing to the resultant opacifying power and brightness. However, paints utilise polymeric binders to fix the pigment and, when in contact with titania, the polymer may oxidise when exposed to sunlight. This effect is known as **chalking** and, in addition to the direct degrading effect of ultraviolet (UV) radiation, is accelerated by the photocatalytic activity of TiO₂, which also is enhanced by the high surface area of this material [25].

The potential for the application of the photocatalytic effect in TiO₂ has attracted considerable interest over the last three decades. Titania photocatalysts are known to be applicable in a range of important technological areas:

- **Energy**
  Electrolysis of water to generate hydrogen [26–30].
  Dye-sensitised solar cells (DSSCs) [31–33].

- **Environment**
  Air purification [34, 35].
  Water treatment [36–40].

- **Built Environment**
  Self-cleaning coatings [34, 38, 39, 41–50].
  Non-spotting glass [47, 51].

- **Biomedicine**
  Self-sterilising coatings [52, 53].

Photocatalytic effect

Photocatalysed reactions for applications such as those mentioned above are facilitated through the presence of adsorbed radicals (from air or water) on the TiO₂ surface [28, 44–47, 49, 54–56]. These radicals, which are atomic species with a free unpaired electron, are formed upon reaction of an adsorbed molecule (such as O₂ or H₂O) with a photo-generated charge carrier (from an electron–hole pair or **exciton**) when TiO₂ is exposed to radiation exceeding its band gap; this radiation normally is in the UV wavelength region (290–380 nm). These electron–hole pairs are formed when an electron is elevated from the valence to the conducting band, leaving behind an electron hole, as shown in Fig. 1.

The electrons in the conduction band facilitate reduction of electron acceptors and the holes facilitate oxidation of electron donors [57]. Examples of the photo-generation of radicals in atmospheric and aqueous environments are given in the following reactions [28, 57–59]:

\[
\text{TiO}_2 + h\nu \leftrightarrow h^+ + e^- \quad (1)
\]

\[
h^+ + \text{H}_2\text{O}_{\text{absorbed}} \leftrightarrow \text{H}^+ + \cdot\text{OH} \quad (2)
\]

\[
h^+ + \text{OH}^-_{\text{absorbed}} \leftrightarrow \cdot\text{OH} \quad (3)
\]

![Fig. 1 Schematic illustration of photo-generation of charge carriers in a photocatalyst](image-url)
\[ e^- + O_2 \text{absorbed} \leftrightarrow {}^*O_2^- \]  

In this notation, an unpaired electron is represented by a point, a valence band electron hole is represented by h\(^+\), and a conduction band electron is represented by e\(^-\).

These mechanisms have been described elsewhere in greater detail for the decomposition of organic pollutants, as shown in Fig. 2 [38, 46, 56, 57, 60] and the splitting of water [26, 27, 29].

The generation of positive and negative charge carriers by UV radiation and their tendency to recombine are competing phenomena, the rates of which govern whether or not a semiconductor can function as a photocatalyst [61]. A key factor in titania’s photocatalytic ability is its high surface area, the same property that contributes to its optical properties. A high surface area leads to a higher density of localised states, which involves electrons with energies between the conduction band and valence band [62]. These electrons are present owing to terminated and unsaturated bonds on the surfaces, and these localised states provide beneficial charge separation in the form of trapping sites for photo-generated charge carriers [63]. Titania has a relatively slow rate of charge carrier recombination in comparison with other semiconductors [64], which is an advantage since it has been suggested that a photo-generated electron–hole pair needs a lifetime of at least 0.1 ns for chemical reactions to be facilitated [64, 65].

In the presence of reactive species adsorbed on the catalyst surface, photogenerated charge carriers may transfer to these adsorbates to form radicals rather than recombine [28]. Thus, an effective photocatalyst also is likely to have a high density of reactive adsorbed species for good performance to be achieved.

Despite the larger experimental band gap of anatase of \(~3.2\) eV, compared with \(~3.0\) eV for rutile [8, 66–68], the photocatalytic performance of anatase generally is considered superior to that of the more stable rutile. This is attributed to a higher density of localised states and consequent surface-adsorbed hydroxyl radicals and slower charge carrier recombination in anatase relative to rutile [58, 60, 69–72], parameters that contribute to improved performance. The higher rate of electron–hole recombination in rutile is considered to result from this material’s typically larger grain size [55, 73] and its resultant lower capacity to adsorb species [60, 74, 75].

It may be noted that, owing to the different crystal structures and associated exposed planes of the two polymorphs, anatase has been reported to have a lower surface enthalpy and lower surface free energy than rutile [76]. Hence, it would be expected that the wetting of anatase by water would be less than that of rutile since higher surface free energies generally contribute to hydrophilicity [77]. Since a high density of adsorbed species would be expected from a hydrophilic material, rutile could be anticipated to exhibit superior photocatalytic performance. It may be noted that there are no reports of rutile’s exhibiting higher levels of adsorbed species.

The photoactivity of anatase and rutile have been examined and interpreted by Sclafani and Herrmann [58] with reference to the densities of surface-adsorbed species. This study showed that higher levels of radicals adsorbed on the anatase surface gives rise to significantly higher photoactivity than rutile. This result was reported to be due to a higher surface area as well as a higher photoactivity per unit of surface area. A similar result was found by Augustynski [71], who reported that surface-bonded peroxo species on anatase were absent from rutile surfaces.

In contrast to the widely reported photocatalytic superiority of anatase, several publications have suggested that, in some cases, rutile may be advantageous for certain applications [23, 25, 78–80]. These studies involved high-surface-area rutile of acicular morphology [25], rutile containing residual anatase [78], and iron-doped rutile [81]. It is possible that electron transfer between rutile and a residual quantity of anatase [78] may facilitate improved photo-oxidative reactions, as in mixed-phase titania catalysts. Therefore, in light of importance of surface area, morphology, and doping, an understanding of the titania polymorphs, their transformation, and the methods by which they can be controlled are likely to be critical to achieving phase-optimised photocatalytic performance.

**Formation and analysis of titania phases**

Phase formation during synthesis of TiO\(_2\)

In the synthesis of TiO\(_2\) films by various methods, the initial crystalline TiO\(_2\) phase formed is generally anatase [82, 83]. From a structural perspective, this could be due to the greater ease of the short-range ordered TiO\(_6\) octahedra in arranging into long-range ordered anatase structure owing to the less-constrained molecular construction of anatase relative to rutile [84]. Alternatively, from a thermodynamic perspective, the more rapid recrystallisation of anatase could be due to the lower surface free energy of this polymorph, despite the lower Gibbs free energy of
rutile [6, 76]. That is, the higher surface free energy of rutile crystallites may favour the crystallisation of anatase.

It should be noted that it is possible to form rutile under near room temperature conditions [54, 83, 103, 104]. Hydrothermal methods of synthesis, which can facilitate the precipitation of crystalline TiO₂ directly from a liquid phase, can be controlled to precipitate rutile. Aside from this method, rutile is obtained only through high-temperature treatment. Table 2 summarises the phases that can result from various synthesis methods at room temperature and at elevated temperatures.

Anatase to rutile transformation

Control of the conditions that affect the kinetics to control the anatase to rutile phase transformation is of considerable interest. This is particularly the case for high-temperature processes and applications, such as gas sensors and porous gas separation membranes [105–107], where the phase transformation may occur, thereby altering the properties and performance of these devices. Therefore, an understanding of the stabilities of the TiO₂ polymorphs, the kinetics of their phase transformation, and the processes involved in controlling them is essential to the ability to obtain single-phase or multiphase microstructures. These issues are critical to the long-term consistency of devices, where retention of anatase or a multiphase microstructure may not be possible, thereby potentially requiring processing designed to produce single-phase rutile. Similarly, limitations in temperature while desiring a specific polymorph, such as rutile, may require manipulation of the materials and processing conditions so as to enhance the direction formation of rutile.

The generation of the phases of TiO₂ depends significantly on the synthesis parameters, which in turn affect the product. The kinetics of these processes typically are considered in terms of temperature and time. In terms of the former, pure bulk anatase is considered widely to begin to transform irreversibly to rutile in air at ~600 °C [6, 66, 99]; however, the reported transition temperatures vary in the range 400–1200 °C [42, 66, 108–112] owing to the use of different methods of determining the transition temperatures, raw materials, and processing methods. The anatase to rutile transformation is not instantaneous; it is time-dependent because it is reconstructive [2, 109, 113]. Consequently, the kinetics of the phase transformation must be interpreted in terms of all of the factors that influence the requisite temperature–time conditions. These parameters for undoped anatase are:

- Particle size
- Particle shape (aspect ratio)
- Surface area
- Atmosphere
- Volume of sample
- Nature of sample container
- Heating rate
- Soaking time
- Impurities (from raw materials and container)
- Measurement technique

The anatase to rutile transition, sometimes referred to as the ART, is a nucleation and growth process [106, 109]. As mentioned, the kinetics of this transition are dependent on variables such as impurities, morphology, sample preparation method, heat flow conditions, etc. In the absence of impurities, dopants, secondary phases, or other types of contamination, rutile forms as fine laths with the product phase’s (100) planes parallel to the (112) planes of the parent anatase [106, 111, 114]. In pure anatase, rutile may nucleate at (112) twin interfaces in anatase [106, 114] as these sites are structurally similar to rutile.

Both anatase, space group I4/amd, and rutile, space group P4₂/mnm, are tetragonal in structure. Both crystal

| Table 2 Common synthesis methods of titanium dioxide and resultant phases |
|----------------|---------------------------------|----------------|----------------|----------------|
| Synthesis method | Mechanism | Phases formed | References |
| Room temperature hydrolysis of TiCl₄ | Precipitation from room temperature solutions of TiCl₄ | ✓ | [85, 86] |
| Room temperature sol–gel synthesis | Hydrolysis of TiCl₄ or an organo-metallic compound | ✓ | [87–90] |
| Flame pyrolysis of TiCl₄ | Combustion of TiCl₄ with oxygen; used in industrial processes | ✓ | [91–93] |
| Solvothermal/ hydrothermal | Precipitation of TiO₂ from aqueous or organic solution at elevated temperatures | ✓ | [66, 84, 94–99] |
| Chemical vapour deposition | Spraying of Ti-bearing solution | ✓ | [100, 101] |
| Physical vapour deposition | Deposition of evaporated Ti and its subsequent oxidation | ✓ | [21, 102] |
structures consist of TiO$_6$ octahedra, sharing four edges in anatase and two in rutile [8, 9, 69, 115, 116]. These structures are illustrated in Figs. 3 and 4.

The anatase to rutile transformation is reconstructive, which means that the transformation involves the breaking and reforming of bonds [117]. This is in contrast to a displacive transformation, in which the original bonds are distorted but retained. The reconstructive anatase to rutile transformation involves a contraction of the $c$-axis and an overall volume contraction of $\sim 8\%$ [109, 118, 119]. This volume contraction explains the higher density of rutile relative to anatase. In the course of the transition to rutile, the (112) planes in anatase are retained as the (100) planes in the rutile product [67, 115]. The $c$-axis of anatase appears to be significantly longer than that of rutile only because anatase has more atoms per unit cell than rutile.

Phase differentiation and quantification of anatase/rutile ratio

**X-ray diffraction**

Examination of the kinetics of the anatase to rutile phase transformation may involve assessment of the relative quantities of anatase and rutile following heating in specific conditions of heating rate, temperature, and time in order to examine the effects of parameters such as dopants, particle size, and atmosphere on the resultant phase assemblage. Quantification of phase proportions usually is carried out by X-ray diffraction (XRD) [1, 78, 81, 105, 120–125]. Such analyses often are done using the method of Spurr and Myers [126], which utilises the ratio of the rutile (110) peak at 27.355° 2$\theta$ to the anatase (101) peak at 25.176° 2$\theta$. The ratio of the intensities of these peaks, $I_R/I_A$, is used in the empirically determined formula in Eq. 5 to give the weight fractions of anatase and rutile:

$$\frac{W_R}{W_A} = 1.22 \frac{I_R}{I_A} - 0.028$$

Despite the number of the above researchers who have utilised this formula in their investigations into the ART, consideration of the relevant issues reveals a number of factors that may affect the accuracy of the results:

- **Preferred orientation**: rutile and/or anatase crystallites may be present in preferred orientation owing to morphological and/or sample preparation effects, which may lead to altered XRD relative peak intensities.
- **Encapsulation**: rutile crystallites may grow as an overlayer of rutile on anatase particles [80] or, alternatively, rutile may form in the bulk of the anatase grains, leaving a surface layer of anatase on rutile particles [112], thereby compromising the basis for the intensities of the XRD peaks.
- **Lattice distortion**: the presence of dissolved dopants and/or impurities, especially if differential solubility occurs, may alter the peak heights and areas, thereby altering the relative intensities of the XRD peaks.
- **Degree of crystallinity**: the presence of dopants may increase (nucleation) or decrease (lattice distortion/stress) the degree of crystallinity, which would alter the consequent peak intensities, particularly if these dopants are preferentially present in one of the phases.
• **Grain size**: the transition to rutile is accompanied by significant grain growth [127, 128], resulting in large rutile grains and small anatase grains, which would alter the ratios of XRD peak intensities.

• **Morphology**: rutile may form in an acicular morphology, which would alter the XRD peak intensities of these grains relative to typically equiaxed rutile [129].

• **Surface nucleation of rutile**: enhanced surface nucleation of rutile owing to heat (thermal gradients) and segregation (chemical gradients) effects would increase the XRD peak intensities of this phase; this would be similar to encapsulation.

Thus, it is important to consider that the validity of the data resulting from Eq. 5 depends on adequate assessment of the preceding parameters and the application of materials and processes designed to achieve samples of representative chemistry, mineralogy, and microstructure. If these cannot be attained, alternative materials, fabrication methods, and analytical techniques may be required.

### Laser Raman microspectroscopy

Laser Raman microspectroscopy also has been employed effectively to differentiate the anatase and rutile [25, 80, 124, 130–133]. The titania polymorphs exhibit distinct Raman spectra that can be used for qualitative and quantitative mineralogical analysis. Laser Raman microspectroscopy has been applied quantitatively for on-line process control in the manufacture of titania powder by Huntsman Tioxide, Inc. [134].

Laser Raman microspectroscopy has several advantages:

- Minimal or no sample preparation
- Nondestructive
- Local and general phase analyses (≤1 μm beam diameter)
- Mapping capability
- Rapid analyses (~1 min scan)
- No preferred orientation effect
- Greater sensitivity than XRD
- Sensitive to nanoscale phases

Figure 5 contrasts experimental scans of identical samples of rutile and anatase TiO₂ by XRD and laser Raman microspectroscopy carried out by the authors and confirmed by comparison with reference patterns and spectra [135–137].

The main strength with laser Raman microspectroscopy is its applicability to nanoscale thin films. When glancing-angle XRD is used, the instrument requires considerable calibration, the peaks are very diffuse, and the background is such that any amorphous phase, if present, cannot be ascertained. With laser Raman microspectroscopy, none of these problems are experienced, so the analyses are more efficient and more sensitive.

### Differential solubility

A more direct method of measuring the rutile/anatase proportions was employed by Ohno et al. [23, 78, 138]. This method involved using hydrofluoric acid (HF) to dissolve the anatase phase, leaving the rutile phase intact. This method has the disadvantages of health risk, potential loss of material during washing, loss of anatase, and potential for slight solubility of rutile. However, this method could be useful for the removal of trace levels of anatase in order to enhance the purity of rutile powders.

### Impedance spectroscopy

Another method of measurement was employed by Suresh et al. [139], who applied impedance spectroscopy to examine the anatase to rutile transformation. Since this method relies on the difference in resistivity between the two phases, it suffers two innate uncertainties associated with electrical resistivity measurement. First, the resistivity is a function of the grain connectivity, which is a microstructural factor rather than a mineralogical one. Second, dopants and impurities have a profound influence on the electrical properties, so their presence inevitably would affect the resistivities in a differential manner. Despite these reservations, this method has been used in conjunction with
XRD and differential thermal analysis (DTA) to ascertain the effect of doping with silica [140].

In general, the XRD method of phase analysis is the most generally used for bulk qualitative and quantitative analyses (using Eq. 5). Experimental calibration has demonstrated that, despite a number of potential flaws in the quantitative method of XRD phase analysis of titania, this method gives reasonably accurate and consistent results [129].

Rutile–anatase mixtures

Mixed-phase photocatalysts with rutile–anatase compositions have been reported to exhibit enhanced photoactivity relative to single-phase titania [23, 34, 54, 55, 68, 73, 80, 110, 112, 125, 130, 131, 141–146]. It is considered widely that this is a result of improved charge carrier separation, possibly through the trapping of electrons in rutile and the consequent reduction in electron–hole recombination [78, 117, 147]. Surface trapping of holes together with lattice trapping of electrons has also been reported [78, 125]. In consequence, Degussa P-25 is marketed as a mixed-phase titania photocatalyst and it is utilised as a reference material in many studies. It should be noted that the high performance observed for Degussa P-25 is reported to stem primarily from this material’s high specific surface area [58]. This nanocrystalline material, formed by flame pyrolysis, consists of 80 wt% anatase and 20 wt% rutile. However, Bacsa and Kiwi [54] reported that a photocatalyst containing 70 wt% anatase and 30 wt% rutile, with a surface area of 72.0 m²/g exhibited greater photocatalytic performance than the Degussa P-25 reference, which has a specific surface area of 49.2 m²/g.

Since the particle size and surface area depend strongly on the preparation method, it would be misleading to interpret the photocatalytic performance solely in terms of the phase composition. As an example of the importance of the effect of the processing conditions on these parameters, variations in grain size, crystallite size, surface area, morphology, and phase distribution obtained during sol–gel synthesis have been observed to be highly dependent on the precursor solutes and solvents used [36, 46, 54], calcining temperatures [55, 110, 148, 149], and pH [54, 73, 83, 114, 139, 150].

Within a certain set of processing protocols, it is possible to obtain mixed-phase photocatalysts of nominally pure titania through variation of the kinetics (heating rate, soak time, and soak temperature). In such cases, a comprehensive understanding of the following effects must be in place:

- Time
- Temperature
- Atmosphere
- Intrinsic chemical composition (purity of raw materials)
- Extrinsic contamination (from processing)
- Chemical homogeneity (e.g., segregation)
- Microstructural homogeneity (e.g., grain boundary precipitates)
- Thermal homogeneity during heating (e.g., resulting from sample size and shape)
- Mineralogical phase assemblage
- Particle size distribution
- Agglomerate size distribution
- Grain morphology
- Agglomerate morphology

In contrast to the processing of nominally pure samples, mixed-phase titania of controlled rutile/anatase ratios also can be obtained by doping. However, doping inevitably affects the semiconducting properties of materials, which may be disadvantageous to the photocatalytic performance. Alternatively, as discussed subsequently, doping also may act advantageously through reduction in the band gap and improvement in the charge carrier separation and associated extended exciton lifetime [18, 68, 73, 151].

Thermodynamics and kinetics of the anatase to rutile phase transformation

Stability of the TiO₂ phases

At all temperatures (Fig. 6) and pressures (Fig. 7), rutile is more stable than anatase. This has been confirmed by thermodynamic studies [5, 9, 76, 148, 152], which show that negative pressures would be required for anatase to be more stable than rutile [2]. The transformation to rutile thus is irreversible.

Table 3 gives the reported standard-state thermodynamic data for the anatase to rutile phase transformation.

![](image-url)

**Fig. 6** Plot of Gibbs free energy of anatase and rutile versus temperature [153]
Over the last 50 years various studies involving the kinetics of the transition to rutile have reported various transition temperatures. A comprehensive survey of these reported temperatures is shown in Table 4.

Table 4 shows that there are different methods that be used to define the phase transformation temperature. As discussed previously, the temperature at which the transition is observed depends on a number of parameters and so it is not surprising that a range of values has been reported. However, assessment and consideration of these data lead to the general conclusion that fine powders of high purity show phase transformation at temperatures from ~600 to 700 °C.

Excluding studies which used titania of unusually small particle size or long soak times, and studies which did not use onset temperatures as the defined transition

Table 3 Summary of thermodynamic data from literature

| Publication year | $\Delta H_{298}$ (kJ/mol) | $\Delta S_{298}$ (J/mol K) | $\Delta G_{298}$ (kJ/mol) | Reference |
|------------------|---------------------------|---------------------------|---------------------------|-----------|
| 1967             | -5.19                     | 0.42                      | -5.32                     | [148]     |
| 1971             | -11.7                     | 0.42                      | -11.84                    | [153]     |
| 2009             | -1.70                     | 0.556                     | -1.87                     | [5]       |

These data indicate that there remains some uncertainty about the energetics of the phase transformation

Table 4 Summary of reported temperatures for the ART

| Year      | Temp. (°C)$^a$ | Fabrication details | Definition | Ref |
|-----------|----------------|---------------------|------------|-----|
| 1961      | 610            | Highly pure powder  | Onset temperature observed by XRD after firing for 24 h | [113] |
| 1965      | (1190)         | Powders from three different suppliers | Apexes of DTA peaks (data not shown) | [109] |
|           | (1138)         |                      |            |     |
|           | (1115)         |                      |            |     |
| 1968      | 610            | Commercially available reagent grade powder | Onset temperature observed by XRD after extended firing (1–5 days) | [154] |
| 1995      | 390            | Sol–gel synthesised powder | Appearance of detectable rutile peak by XRD after 1 week | [123] |
| 1996      | 675            | Sol–gel synthesised powder | Appearance of detectable rutile peak by XRD after 4 min | [1] |
| 1997      | (787)          | Sol–gel synthesised powder | Reported value: Apex of DTA peak | [155] |
|           | 720            |                      | Corrected value: Onset of DTA peak |     |
| 1997      | 465            | 4–6 nm particles prepared through a sol–gel method | Appearance of detectable rutile peak by XRD | [156] |
| 1999      | (616)          | Sol–gel synthesised powder | Reported value: 50% transformation observed by XRD | [157] |
| 2001      | (680)          | Sol–gel synthesised powder | Reported value: 50% transformation observed by XRD | [82] |
|           | 600            |                      | Onset temperature from graph |     |
| 2001      | 600            | Sol–gel synthesised powder | Appearance of detectable rutile peak by XRD | [108] |
| 2002      | (700)          | Sol–gel synthesised powder | 50% transformation observed by XRD | [124] |
|           | 600            |                      | Appearance of detectable rutile peak by XRD |     |
| 2005      | 600–700        | Highly pure nanocrystals synthesised from$\text{TiCl}_4$ Sol–Gel | Appearance of detectable rutile peak by XRD | [99] |
| 2007      | (900)          | Sol–gel synthesised powder | Reported value: apex of broad DTA peak (data not shown) | [158] |

$^a$ Data in parenthesis do not correspond to the onset of the transformation
temperatures, it can be seen that the reported transition onset temperatures, as determined by XRD, appear to converge around 600°C. Although it is difficult to ascertain the intrinsic behaviour of titania, this is accepted to be the region of the onset temperature of the anatase to rutile transformation in bulk pure anatase in air [2, 7, 123, 154]. This can be seen in Fig. 8.

The one early study from 1965 [109] that reported anomalously high temperatures may have obtained these results owing to the use of powders contaminated with impurities that inhibit the phase transformation, as discussed subsequently, the use of the differential thermal analysis (DTA) peak apices rather than the onsets, and/or the use of very high heating rates.

The data in Table 4 suggest that DTA analysis yields higher transformation temperatures than XRD analysis. This probably is a result of the sigmoidal form of the kinetics curve, as shown subsequently. That is, the phase transformation commences slowly at an onset temperature of ~600°C and so it is more likely that the exotherm maximum (viz., the apex) is observed later in the phase transformation process, when the transformation rate is greatest; the rate of transformation subsequently decreases along with the corresponding latent heat as anatase converts to rutile. Despite the potential uncertainties, DTA peak apices can be considered to be applicable for comparative analysis of the phase transformation.

The assessment of DTA data must be done with care since the curves and their interpretation can be influenced significantly by experimental conditions that can cause shifts to lower temperatures: (1) a slow heating rate results in peak broadening, which lowers the onset temperature; (2) a slow heating rate also causes the entire peak to shift; (3) fine particle size shifts and narrows the entire peak; and (4) the static enclosed atmosphere induces a slightly reducing atmosphere, which enhances oxygen vacancy formation in oxides.

Rao [113] carefully examined the kinetics of the phase transformation, based on a soak of 24 h and XRD analysis, using very pure powders and fit the data to an Arrhenius relation. The rate of transformation decreased with temperature to a practical limit 610 ± 10°C, at which point the transformation became immeasurably slow. Despite the age of this study, it is consistent with later studies [154, 157, 159, 160].

Since rutile is the equilibrium phase [2, 7, 9, 20, 76, 113, 148, 152], the presence of anatase demonstrates that these studies cannot represent equilibrium conditions, which normally employ phase diagrams for illustration. Dachille et al. [154] reported what was considered to be an pressure–temperature diagram approximating equilibrium for TiO2, which is shown in Fig. 8. The key observation of these data is the apparent anatase to rutile phase transformation conditions of 605°C at 1 atm (101 kPa) pressure, which is quite similar to the values indicated in Table 4.

Jamieson and Olinger [2] subsequently qualified the description of this diagram by describing the phase boundaries as reaction boundaries. Current practice is to refer to functional diagrams such as these as behavioural diagrams.

Although it is widely accepted that rutile cannot be transformed to anatase. It has been suggested that at high pressures rutile can transform to the $alpha$-PbO2 structured TiO2 II polymorph [12, 161]. This behaviour is outlined by the inferred behavioural diagram shown in Fig. 9, which is derived from mineralogical samples exposed to high pressures in the earth’s crust.

Kinetics of the anatase to rutile phase transformation

For the transformation of anatase to rutile to proceed at a measurable rate, sufficient thermal energy is required to facilitate the rearrangement of atoms. As described above, it is likely that, for typical bulk titania powders (i.e., not nanoparticles), this energy requirement is reached at ~600–700°C in air in the absence of dopants or impurities,
although this is subjective since impurities are always present at a finite level. As will be discussed in detail subsequently, the transformation can be enhanced or impeded by influencing the rearrangement of the atoms in the anatase and rutile lattices. It is perceived widely that the most important factor affecting the phase transformation is the presence and amount of defects on the oxygen sublattice, viz., TiO$_2$$_{-x}$ [67, 157, 162, 163]. Ease of rearrangement and transformation are enhanced by relaxation (lessening of structural rigidity) of the large oxygen sublattice through the increased presence of oxygen vacancies [109, 164, 165].

This effect has been shown through firing in different atmospheres, where neutral or reducing conditions of low oxygen partial pressure generally greatly enhance the anatase to rutile transformation [3, 67, 166], although contrary effects have been reported as vacuum conditions have been observed to slow the phase transformation [109]. This could be a result of reduced heat transfer because vacuum conditions give lower convective heat transfer than air. The promotion of the phase transformation through the use of a reducing atmosphere is considered to be due largely to the increased levels of oxygen vacancies during heating in such atmospheres.

Figure 10 shows the effects of different experimental conditions on the kinetics of the anatase to rutile transformation in four different samples. Although these data are limited, the interpretations of the trends are straightforward:

Sample a: Doping with Fe$_2$O$_3$ (assuming substitution of Ti$^{4+}$ by Fe$^{3+}$) in a reducing atmosphere can increase the levels of oxygen vacancies by three potential mechanisms: (1) the maintenance of charge balance; (2) spontaneous reduction Fe$_2$O$_3$ to Fe$_3$O$_4$ or FeO, which are thermodynamically stable at temperatures as low as 400 °C and low oxygen partial pressures [168]; and (3) reduction of TiO$_2$ to TiO$_2$$^{-x}$

Samples b and d: Assuming that powders used in these two studies are similar, the only difference is the temperature. If this comparison is valid, then it is clear that the phase transformation is accelerated with the use of the higher temperature, which is as expected.

Samples c and d: Doping with MnO$_2$, while suggesting that no effect on oxygen vacancy formation should be observed, actually is similar to Fe$_2$O$_3$ because Mn$^{4+}$ reduces spontaneously in air to Mn$^{3+}$ and then Mn$^{2+}$, with the transformation’s near-completion by $\sim$450 °C [169]. Again, the assumption of substitution of Ti$^{4+}$ by Mn$^{3+}$ and Mn$^{2+}$ requires the generation of oxygen vacancies.

**Effects of impurities and dopants on the anatase to rutile phase transformation**

**Dopant effects**

Many studies of photocatalysis by TiO$_2$ have attempted to utilise dopants in order to improve the photocatalytic activity. In general, the role of dopants is considered to be:

- Reduction of the band gap in titania [18, 151]
- Introduction of mid-gap states [56, 163]
- Improvement in charge carrier separation [68, 73]
- Increase in the levels of surface-adsorbed species (e.g., hydroxyl radicals) [43]

It often is not appreciated that all titania is contaminated with some levels of impurities. The presence of unintentional impurities or intentional dopants has a strong effect on the kinetics of the anatase to rutile transition [170]. Variable results have been reported in the sense that dopants can have the effect of hindering or enhancing the transition to rutile. In the case of substitutional solid solution formation, dopant ions can enter the anatase lattice and influence the level of oxygen vacancies, thereby promoting or inhibiting the transformation. In the case of interstitial solid solution formation, lattice constraint may result in destabilisation or stabilisation, depending on size, valence, and content effects, again promoting or inhibiting the transformation.
If the solubility limit for impurities or dopants is exceeded, then their precipitation can facilitate the phase transformation through heterogeneous nucleation [67, 171].

Cationic dopants

Numerous cationic dopants have been investigated in terms of their effect on the kinetics of the anatase to rutile transition. It has been suggested that cations of small radii and low valence accelerate the transition to rutile owing to the increase in oxygen vacancies that result from the assumed substitution of Ti$^{4+}$ ions with cations of lower valences [43, 109, 157, 164, 165]. Charge neutrality thus requires an increase in the level of oxygen vacancies and/or the formation of Ti interstitials of lower valence [82, 172]. Conversely, when cations of valence higher than 4 are assumed to substitute for Ti ions on the anatase lattice, this gives rise to the annihilation of existing oxygen vacancies and the formation of Ti interstitials of the same or lower valence. These processes can be viewed in light of the inertia to alteration (through ionic transport) of the relatively large and rigid oxygen sublattice, which largely determines the structural stability and the capacity to reorganise the chemical bonds to form rutile. From these considerations, the assumption of substitutional solid solubility leads to the conclusion that small cations of low valence (<4) should promote the anatase to rutile transformation and large cations of high valence (>4) should inhibit it.

However, the assumption of substitutional solid solubility may be incorrect and interstitial solid solubility occurs. In this case, the insertion of a cation results in constraint of the required lattice contraction largely in the c direction upon the transformation from anatase to rutile [173], without apparent effect on the charge neutrality. Although there are reports of interstitial stabilisation of the titania lattice and consequent inhibition of the transformation [155, 174, 175], there do not appear to be any reports of destabilisation (from structural instability) and consequent promotion of the transformation.

A comprehensive summary of the cationic dopant effects from the literature sources is given in Table 5.

The effects of the dopant cations in terms of the valences and ionic radii are illustrated in Fig. 11. These data have been compiled on the basis of the following conditions:

- Shannon-Prewitt ionic radii in sixfold coordination [173], which is the case for both anatase and rutile [14], were used.
- The most common valence for each cation has been used, without regard to oxidation–reduction effects (except as noted below).
- The valences for Mn, Fe, and Co are assumed to be 2 as these species would be likely to reduce spontaneously to show this valence state upon heating at temperatures approaching the ART temperature [167, 169, 192].
- Inhibition or promotion of the phase transformation is reported on the basis of the effects described in the literature sources given in Table 5.

### Table 5 List of dopants in titania

| Phase transformation inhibitors | Cation | Dopant phases used |
|---------------------------------|--------|-------------------|
| Al                              | AlOOH [176], Al(OCH₃H₃)₃ [108], AlCl₃ [119, 177], Al(NO₃)₃ [175] |
| Au                              | HAuCl₄ [178] |
| B                               | BCl₃ [120] |
| Ba                              | Ba(NO₃)₂ [157] |
| Ca                              | Ca(NO₃)₂ [157] |
| Ce                              | CeO₂ [179] |
| Dy                              | Dy₂O₃ [180] |
| Eu                              | EuCl₃ [70], Eu₂O₃ [56] |
| Er                              | Er(NO₃)₃ [157], Er₂O₃ [181] |
| Fe                              | FeCl₂ [171] |
| Ho                              | Ho₂O₃ [181] |
| La                              | La(NO₃)₃ [157, 182], La₂O₃ [181] |
| Mn                              | Mn(NO₃)₂ [183], <1 mol% Mn(CH₃COO)₂ [184] |
| Nb                              | NbCl₅ [185] |
| Nd                              | Nd₂O₃ [181] |
| P                               | PO₄H₂K [118], POCl₃ [120] |
| Si                              | Si(OCH₃H₃)₃ [82, 108, 128, 175], SiCl₄ (g) [120] |
| Sm                              | SmCl₃ [70], Sm₂O₃ [181] |
| Sr                              | Sr(NO₃)₂ [157] |
| Tb                              | Tb₂O₃ [181] |
| Tm                              | Tm₂O₃ [181] |
| Y                               | Y (NO₃)₂ [157], Y₂O₃ [181] |
| Zr                              | Zr(OCH₃H₃)₆ [108, 186], Zr(SO₄)₂ [66], ZrOCl₂ [87] |

| Phase transformation promoters | Cation | Dopant phases used |
|--------------------------------|--------|-------------------|
| Al                              | AlCl₃(g) [187] |
| Cd                              | CdO [165] |
| Co                              | CoO [165] |
| Cr                              | CrCl₃ [185] |
| Cu                              | CuO [109, 127, 164] |
| Fe                              | Fe [41], Fe₂O₃ [127, 164, 167] |
| Li                              | LiF [118] |
| Mn                              | MnO₂ [164, 165], Mn(NO₃)₂ [183, 188], MnSO₄ [67] |
| Na                              | NaF [165] |
| Ni                              | Ni(NO₃)₂ [128], Ni(CH₃COO)₂ [189] |
| Sb                              | Sb₂O₃ [190] |
| Sn                              | SnO₂ [158], SnCl₂ [149] |
| V                               | VO₂ [128], V₂O₅ [191] |
| Zn                              | ZnO [165] |
In the case of cations for which contradictory effects are reported (Mn and Fe), the most common finding was used.

The formula of the boundary line is

\[ y = -0.0455x + 0.2045 \]

where \( x \) is the valence and \( y \) is the ionic radius.

The data shown in Fig. 11 should be considered indicative owing to the potential presence of impurities, the vagaries in the reported experimental procedures, and the assumptions stated above. For example, in 1960, Iida and Ozaki [127] reported findings which contrary to those of many other studies in that sodium was observed to inhibit the transformation. This seems likely to be due to the use of sodium in the form of a salt which only decomposed at 900 °C.

Anionic dopants

Anions doping is of interest owing to its potential to improve the photocatalytic performance of TiO\(_2\) [18]. There is limited discussion in the literature regarding the effects of foreign anions on the anatase to rutile transformation, probably owing to the uncertainty of whether doping actually occurs. It is reasonable to assume that doping by anions results in the filling of oxygen vacancies. The inhibition or promotion of the phase transformation is likely to depend on size and charge effects, which are as follows:

\[ \text{Cl}^{1-} > \text{N}^{3-} > \text{O}^{2-} > \text{F}^{1-} \]

Nitrogen doping

Size effects: according to Hume-Rothery [193], metal ion dopants whose ionic radii are within 15% of that of the host lattice, which typically is closest packed, can show appreciable solubility. It is assumed that the oxygen sublattice, which also is closest packing, can be considered similar to that of most metals. Unfortunately, the only comparable ionic radii for the two anions is for fourfold coordination [S–P], but the ionic radii of anions vary little across the different negative valences. Since nitrogen is only \( \sim 6\% \) larger than oxygen, then substitution of nitrogen on the oxygen sublattice is possible. Hence, nitrogen is unlikely to destabilise the lattice and so would be expected to be an inhibitor of the phase transformation.

Charge effects: if \( \text{N}^{3-} \) substitutes for \( \text{O}^{2-} \), then the impossibility of establishing charge balance through the formation of \( \text{Ti}^{5+} \) means that oxygen must be removed from the lattice. Since the inclusion of two nitrogen ions is balanced by the removal of three oxygen ions, then the level of oxygen vacancies increases, the lattice can be expected to be destabilised, and the phase transformation is promoted. Accordingly, it has been reported that nitrogen doping increases the level of oxygen vacancies in anatase [117, 162] although there do not appear to be any reports on the effect of chemical nitrogen (rather than physical nitrogen, as in the furnace atmosphere) on the phase transformation.

Fluorine doping

Size effects: fluorine in both fourfold and sixfold coordinations is \( \sim 6\% \) smaller than oxygen, then fluorine replacement on the oxygen sublattice is possible. Hence, fluorine would be expected to be an inhibitor.

Charge effects: if \( \text{F}^{1-} \) substitutes for \( \text{O}^{2-} \), then anion charge balance requires the inclusion of two fluorine ions for each oxygen. This is extremely unlikely from size considerations. The other alternative is the inclusion of one fluorine ion in an oxygen vacancy site and the resultant reduction of the associated \( \text{Ti}^{4+} \) or \( \text{Ti}^{3+} \), which would remain on the cation sublattice; the insertion of fluorine would decrease the level of oxygen vacancies. Further, the expansion of the titanium ionic radius upon reduction [s–p] would cause lattice constraint. Both of these mechanisms would be expected to inhibit the phase transformation. In agreement, fluorine has been reported to inhibit the phase transformation [64, 90].

The annihilation of an oxygen vacancy by fluorine doping is described by Eq. 7:

\[ \text{F}^- + 2\text{Ti}^{3+} + 3\text{O}^{2-} + \Box_a \rightarrow \text{Ti}^{4+} + \text{Ti}^{3+} + 3\text{O}^{2-} + \text{F}^- \]  

(7)

Here, \( \Box_a \) denotes an anion vacancy.
Chlorine doping

Size effects: chlorine in sixfold coordination is \(\approx 33\%\) larger than oxygen. Hence, it is very unlikely to enter (solid state) or be retained in (solution) the titania lattice.

Charge effects: in light of the above, the inclusion of two chlorine ions for one oxygen ion is extremely unlikely. For the same reason, inclusion of one chlorine and titanium reduction also is unlikely. The preceding being said, chlorine has been reported to inhibit the phase transformation [64, 90]. Conversely, atmospheric chlorine also has been reported to promote the phase transformation [3].

Valence changes

An important issue to consider is that certain cationic dopants may exhibit more than one valence and the associated potential for reduction–oxidation reactions [165, 169]. These can result in an increase in the level of oxygen vacancies (promotion of the phase transformation through increase in lattice relaxation) and/or the formation of Ti\(^{3+}\) interstitial (inhibition of the phase transformation through lattice constraint). The fact that Al\(^{3+}\) has only a single valence may explain why it inhibits the phase transformation as opposed to Fe\(^{3+}\), which promotes the phase transformation, although being of similar size and valence to Al\(^{3+}\).

This can be attributed to its likely valence change during heating.

Iron can undergo reduction creating an oxygen vacancy by the reaction in Eq. 8.

\[
2\text{Fe}^{3+} + \text{O}^2^- \rightarrow 2\text{Fe}^{2+} + \square a + \frac{1}{2}\text{O}_2
\]  

(8)

Here, \(\square a\) denotes an anion vacancy.

In the TiO\(_2\) lattice the anatase to rutile phase transformation will be promoted by the formation of oxygen vacancies by the reduction of dopant iron 3+ following the reaction shown in Eq. 9:

\[
\text{Fe}^{3+} + \text{Ti}^{4+} + 2\text{O}^2^- \rightarrow (\text{Fe}^{2+} + \text{O}^2^- + \square a)
+ \text{Ti}_{\text{interstitial}}^{3+} + \frac{1}{2}\text{O}_2
\]  

(9)

The anion vacancy, \(\square a\), enhances the phase transformation.

In the case of aluminium doping of TiO\(_2\), the reaction is

\[
\text{Al}^{3+} + \text{Ti}^{4+} + 2\text{O}^2^- \rightarrow (\text{Al}^{3+} + \frac{3}{2}\text{O}^2^- + \frac{1}{2}\square a)
+ \text{Ti}_{\text{interstitial}}^{3+} + \frac{1}{2}\text{O}_2
\]  

(10)

In the case of doping of TiO\(_2\) by Mn and its reduction from Mn\(^{4+}\) to Mn\(^{2+}\) which occurs spontaneously at temperatures above 400 °C [169] the reaction is described by the following:

\[
\text{Mn}^{4+} + \text{Ti}^{4+} + 2\text{O}^2^- \rightarrow (\text{Mn}^{2+} + \text{O}^2^- + \square a)
+ \text{Ti}_{\text{interstitial}}^{4+} + \frac{1}{2}\text{O}_2
\]  

(11)

The creation of oxygen vacancies in TiO\(_2\) through valence change in Fe was confirmed by Heald and Weiss [167] and Genarri and Pasquevich [192] who used haematite in solid state form as a dopant. In another study, after Fe was added in a 3+ oxidation state form and the doped TiO\(_2\) was heated in air, electron paramagnetic resistance (EPR) showed that Fe was present in the 2+ oxidation state and that this promoted the phase transformation [194].

In contrast, the addition of Fe in the 2 valence, using FeCl\(_2\) and firing in air, resulted in inhibition of the phase transformation [171]. Mössbauer spectroscopy revealed that iron was present interstitially in the 3 valence. If the 2 valence was retained during heating, then size and charge considerations support the view that Fe\(^{2+}\) would be unlikely to substitute for Ti\(^{4+}\). Further, the oxidation of Fe\(^{2+}\) to Fe\(^{3+}\) would require the annihilation of oxygen vacancies, in reverse of reaction 8. Hence, both of these phenomena would be expected to inhibit the phase transformation.

The preceding different outcomes from doping with the same metal highlight the importance of differentiating between the assumption of substitutional and interstitial solid solubility. Again, the same dopant in substitutional form may increase the oxygen vacancy level and promote the phase transition while, in the interstitial form, may enhance lattice constraint and inhibit the phase transformation. This applies to all dopants, although the transition metals of variable valence (e.g., Mn, Fe, Ni), are most likely to be those that show mixed effects.

In contrast, the absence of the likelihood of valence change is more straightforward. In the case of substitution of Ti\(^{4+}\) by Al\(^{3+}\), half an oxygen vacancy and a Ti\(^{3+}\) interstitial are created (Eq. 7), thereby promoting and inhibiting the phase transformation, respectively. Since, the inhibition of the phase transformation by aluminium doping is well known [155, 175, 195], if substitution is the mechanism, then it can be concluded that the effect of lattice constraint is greater than that of lattice relaxation. However, if Al\(^{3+}\) enters the lattice interstitially, then only lattice constraint results, which inhibits the phase transformation and so no information about the relative importance of the two mechanisms is provided.

Predictive analysis

Using data from the literature, it is in principle possible to make a subjective estimate of the potential effects of
dopants for which no data are available. The four deductive considerations used to predict the effects of dopants on the anatase to rutile transition are:

- Most common valence and ionic radii of substitutional cationic dopants in conjunction with Eq. 6 and the equivalent interpolated boundary in Fig. 11.
- The potential for valence changes in dopant cations (e.g., Fe$^{3+}$ → Fe$^{2+}$).
- The potential for oxygen vacancy formation through reduction (e.g., C- and N-doping).
- The known effects of anionic dopants (e.g., F$^-$ and Cl$^-$).

A summary of the reported and predicted effects of dopants on the phase transformation is given as follows.

**Inhibitors**

See Fig. 12.

**Promoters**

See Fig. 13.
Carbon doping

Carbon is an attractive dopant for titanium dioxide photocatalysts as it has been reported to reduce the band gap and improve photocatalytic performance in anatase [20]. Carbon 4+ has an ionic radius of 0.03 nm, which places it close to the line of Eq. 6, making a prediction of its effects on the anatase to rutile transformation based on valence/size considerations uncertain. Moreover, there is an apparent absence of reported data regarding the effects of carbon on the transformation of anatase to rutile. This is a result of the likelihood of carbon oxidation at temperatures below the anatase to rutile phase transformation temperature. However, carbon is a very strong reducing agent and, when retained during firing in an inert atmosphere, it would be likely to enhance the transformation to rutile through the formation of oxygen vacancies. Further, the oxygen-deficient atmosphere created through the use of an inert gas overpressure also would enhance the transformation to rutile through defect formation. In the extreme case, a stable carbide could form by reaction. Effectively, all of these are reduction reactions that take place to different degrees, as indicated:

\[
\text{Defect Formation: } \text{TiO}_2 + x\text{C} \rightarrow \text{TiO}_{2-x} + x\text{CO} \\
\text{Suboxide Formation: } \text{TiO}_2 + \text{C} \rightarrow \text{TiO} + \text{CO} \\
\text{Carbide Formation: } \text{TiO}_2 + 2\text{C} \rightarrow \text{TiC} + \text{CO}_2
\]

Consequently, carbon is predicted to promote the phase transformation, as shown in Fig. 13.

It has been reported that nitrogen doping increases the levels of oxygen vacancies in anatase [117, 162], so it is likely to promote the transition to rutile. However, this has not been reported. In light of the preceding, nitrogen also is predicted promote the phase transformation in Fig. 13.

Importance of doping methods

The preparation technique of doped anatase is critically important because it impacts on the degree of equilibration achieved. There are three general methods by which dopants can be combined with anatase: point contact, surface contact, and molecular level mixing. These are listed below in order of decreasing diffusion distance required for the dopant ions in order to enter the anatase lattice:

- **Dry mixing:** this involves the blending of dry powders of anatase and dopant-bearing phases, such as oxides. Both large particle sizes and inhomogeneous mixing are associated with increased diffusion distances.
- **Wet impregnation:** this method involves mixing dry anatase powder with a dopant-bearing solution, such as dissolved salts or metal-alkoxides [67, 188].
- **Molecular-level mixing:** this method offers the most intimate level of association and involves mixing of a soluble titanium-bearing compound, typically an organometallic, such as titanium isopropoxide, with a soluble dopant-bearing compounds in an organic or aqueous solution. This level of mixing often is obtained through the use of doped sol–gels or co-precipitation.

Doping methods that involve larger diffusion distances for the dopant compounds to enter the titania lattice may diminish the inhibiting or promoting effect of the dopant on the anatase to rutile phase transformation since this may take place before the dopant has entered the anatase lattice.

Oxide dopants

Alumina, silica, and zirconia and have been used to stabilise anatase [66, 108, 120, 174, 196]. It has been suggested that Al, Si, and Zr stabilise anatase by occupying interstices, thereby distorting the anatase lattice and restricting the lattice contraction involved in the transformation to rutile [120]. Conversely, Yang and Ferreira [175] have suggested that the observed contraction in lattice parameters upon SiO₂ and/or Al₂O₃ doping is evidence of solid solubility. Another study of the effects of SiO₂ doping on the lattice parameter of anatase [82] also suggested that Si⁴⁺ enters substitutionally, thereby decreasing the lattice parameter of anatase (and forming interstitial Ti⁴⁺). Further, it is possible that the distortion of the lattice by the doping restricts the ionic rearrangement similar to that of interstitial ions. Also, the presence of undissolved SiO₂, possibly as a grain boundary glassy phase, has been suggested to inhibit diffusion and reduce anatase interparticle contact, thus reducing the number of available heterogeneous nucleation sites [82].

The use of solid-state dopants introduces the scope for heterogeneous rutile nucleation at the surfaces of the dopant material. This may be the case for tin oxide which has been suggested to facilitate epitaxial growth of rutile due to its similar lattice parameters [158, 197]. Such rutile nucleation has been reported to accelerate the transformation to rutile, causing it to take place rapidly so that coarsening does not have the time to occur. This study supports the view that the grain growth typically associated with the phase transformation is not a cause but an effect.

Other considerations

Effects of firing atmosphere on the anatase to rutile transformation

Rutile is reported widely to exhibit oxygen deficiency and can be described more appropriately as having the formula
TiO$_{2-x}$ [163, 198–200]. Such a stoichiometry requires, in principle, the presence of titanium lattice ions, unintentional impurities, and/or intentionally added dopants of valences lower than 4 in order to maintain charge balance. This nonstoichiometry may be present in anatase as well, although this appears not to have been discussed in the literature. The oxygen vacancies in anatase can be expected to enhance the transformation to rutile owing to the facilitated rearrangement of ions.

In contrast to the use of dopants, the atmosphere used during heating of anatase may affect the probability and kinetics of the transformation to rutile. That is, inert (viz., noble gases) or reducing atmospheres (viz., hydrogen) can be expected to increase the number of oxygen vacancies in the anatase lattice (relative to heating in air), thereby promoting the transformation to rutile. Conversely, heating in air or O$_2$ can be expected to inhibit the transformation owing to the filling of vacancies. These effects have been observed in various reports [67, 109, 167, 192], typically through reporting the transformation temperature.

Formation of crystalline TiO$_2$ from solutions

Alternative methods of obtaining rutile, anatase, or brookite directly from solution at relatively low temperatures have been investigated using solvothermal processes [83, 94–96, 121, 197, 201–203]. These approaches involve the use of titanium-bearing solutions of varying pH. When titanium alkoxides undergo hydrolysis, the result is agglomerated titanium hydroxide groups which are often in the form of white amorphous precipitates [46, 95, 204]. Such solutions also can be synthesised using TiCl$_4$ [96, 99, 103, 104, 202] which can result in crystalline phases of high purity. The use of a low pH limits condensation reactions [197], imposes a repulsive charge and limits the size of these precipitates resulting in the presence of free [Ti(OH)$_x$(OH$_2$)$_{6-x}$]$_{4-x}^{(4-x)+}$ octahedra in solution [197, 201], possibly also through dissolution of amorphous particles [201]. These octahedra can then link together to give crystalline TiO$_2$ phases. Higher pH levels are more conducive to the formation of amorphous titania as octahedra are not free in solution and therefore less able to link together to form a crystalline phase.

Upon heating, the manner in which [Ti(OH)$_x$(OH$_2$)$_{6-x}$]$_{4-x}^{(4-x)+}$ octahedra join determines the crystalline phase that precipitates. If the octahedra have sufficient time to join linearly, with two edges of each octahedron’s being shared (see Figs. 3, 4), the free energy is lowered resulting in the formation of the equilibrium phase rutile in the form of grains elongated in the [001] direction [201, 202]. Alternatively, if the crystallisation rate from the solution is increased, the octahedra may tend to join at right angles as there are more positions for linkage available in this manner, such crystallisation would result in anatase formation which is statistically favoured owing to the more flexible zigzag construction of this polymorph, with four edges of each octahedron’s being shared (see Figs. 3, 4) [201].

Slower crystallisation will yield the rutile phase and more rapid crystallisation favours anatase. This reasoning may also explain why anatase is generally the product phase from most industrial titania manufacturing methods which tend to involve rapid crystallisation of titania through flame pyrolysis of TiCl$_4$.

Morphological effects

The grain morphology plays an important role in photocatalytic applications of titania. Nanocrystallinity has been shown to lower the material’s densification temperature and enhance its photocatalytic activity [73, 76, 110, 205]. A common goal is to inhibit the grain growth of titania during heating as this gives a higher surface area and thus improved performance.

Bulk rutile is more stable thermodynamically than anatase at all temperatures and pressures (Figs. 7, 8) owing to its lower free energy [73, 76, 105, 110, 206]. However, the lower surface energy of the anatase planes relative to those of rutile [76] cause the former to be more stable for crystallites of extremely small sizes and correspondingly high surface areas. In these cases, surface energy considerations outweigh those of bulk thermodynamics and so, for crystallites below a critical size (45 nm [170], 14 nm [76], and 11 nm [207]), anatase has a lower total (bulk and surface) free energy [73, 76, 206]. Further, the size above which rutile becomes more stable depends on stresses [76].

Despite this, it has been reported that larger anatase grains (i.e., predominantly bulk thermodynamics) transform to rutile more slowly than finer grains (i.e., predominantly surface thermodynamics). This probably is due to the lower surface energy and fewer interfaces at which rutile can nucleate. Thus, the transition to rutile and grain growth of anatase can be considered as competing phenomena [205].

Significant rutile grain growth is exhibited as the anatase to rutile transition proceeds [67, 73, 108, 158, 170, 176, 177, 208]. Rutile grains coarsen at the expense of neighbouring anatase during coalescence until the large rutile grains begin to impinge on each other [111, 114]. This increase in grain size causes a decrease in surface area and a consequent decrease in photocatalytic activity [55, 73, 131].

Grain boundary effects

Dopant oxides that do not dissolve and remain on the grain boundaries restrict the grain growth and reduce...
interparticle contact, thereby impeding the phase transformation [87, 170].

Also, the presence of a liquid or vitreous phase on the grain boundaries generally represents a hindrance to diffusion and so it is expected that titania contaminated with silica or another glass former would be likely to inhibit the phase transformation [209].

Summary

An understanding of the transformation of anatase to rutile is of great importance to those studying TiO2 for photocatalysis or other applications. The phase composition of the material has significant consequences on its properties and performance and therefore it may be desirable to enhance or inhibit the transformation to give a particular phase or phase mixture subsequent to thermal treatment.

At all temperatures and pressures, rutile is the stable phase of TiO2. Anatase is metastable but it can be considered to be kinetically stabilised at lower temperatures. Although rutile is the more stable phase from a thermodynamic point of view, anatase frequently is the product phase in the synthesis of TiO2 owing to its less constrained structure and consequent enhanced kinetics of formation. The phase transformation of kinetically stabilised anatase to thermodynamically stable rutile does not have a distinct transformation temperature. Rather, it shows a gradual onset, the temperature of which is highly dependent on numerous parameters pertaining to the nature of the material, the thermal treatment, and measurement method. The transformation rate to rutile increases exponentially as the temperature increases.

The transformation of anatase to rutile can be promoted, accelerating its kinetics and causing it to become observable at lower temperatures and to progress more quickly. Alternatively, the transformation can be inhibited, retarding the kinetics and causing it to become observable only at higher temperatures. The latter also can be referred to as the stabilisation of anatase even though it is the less stable phase.

The kinetics of the anatase to rutile transformation are affected strongly by oxygen defect levels, where oxygen vacancies enhance the transformation. The oxygen defect levels are influenced by atmospheric conditions, reduction or oxidation reactions, unintentional impurities, and intentional dopants.

Dopants are an effective method for influencing the kinetics of the anatase to rutile transformation. Various cationic dopants have been shown to have a strong influence on the phase transformation through the change in oxygen vacancies. This effect can be seen clearly in Fig. 11, which separates inhibitors and promoters in terms of the combined effects of their ionic radii and valence. Substitutional cationic dopants can increase (promoters) or decrease (inhibitors) oxygen vacancy levels through valence effects, alteration in cation coordination, or reduction/oxidation effects. However, the presence of interstitial ions (lattice constraint) and the reduction of interparticle contact (grain boundary phases) in anatase act to inhibit the transformation. Most doping ultimately increases the thermal stability of anatase. The effects of anionic dopants have been reported to a lesser degree.

The consideration of the mechanisms by which the anatase to rutile transformation is inhibited or promoted can enable prediction of the effects that various dopants and experimental conditions are likely to have on the chemistry, mineralogy, and phase assemblage of the resultant titania, as shown in Figs. 12 and 13. This ultimately can assist in facilitating the optimisation of the activity of titanium dioxide photocatalysts.

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