Mechanisms of fault mirror formation in carbonate rocks

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Abstract

The development of smooth, mirror-like surfaces provides insight into the mechanical behaviour of crustal faults during the seismic cycle. To determine the physico-chemical mechanisms of fault mirror formation, we investigated carbonate fault systems in seismically active areas of central Greece. Using multi-scale electron microscopy combined with Raman and electron energy loss spectroscopy we show that fault mirror surfaces do not need to develop from nanogranular volumes, as suggested by previous investigations. Instead, we demonstrate that decarbonation is the main transformation process that leads to the formation of smooth surface coatings in these faults. Piercement structures on top of the fault surface indicate calcite decarbonation, producing CO\textsubscript{2} and lime (CaO), where lime subsequently reacts to portlandite (Ca(OH)\textsubscript{2}) under hydrous conditions. Nanoscale imaging and electron diffraction reveals a thin coating of a non-crystalline material sporadically mixed with nano-clay, forming a complex-composite material that smooths the slip surface. Spectroscopic analyses reveal that a thin layer of non-crystalline carbon covers
the slip surface. We suggest that, during the post-seismic period, ordering (hybridisation) of amorphous carbon led to the formation of partly-hybridised amorphous carbon but did not reach full graphitisation. Calcite nanograins, < 50 nm in size, are spatially associated with the carbon and indicate that the decomposition products acted as a crystallisation medium. Within this medium portlandite back-reacted with CO$_2$ to form nanocrystalline calcite. Hence, we suggest that the nanograins are not the result of comminution during slip but originate from pseudomorphic replacement of calcite after portlandite. Our results suggest that calcite decarbonation products may develop across the entire fault surface, controlling the formation of carbonate fault mirrors, and may facilitate slip on a decarbonation-product glide film.

1 Introduction

Brittle deformation of upper-crustal rocks can result in high-magnitude seismic events (Scholz, 1998). Mirror slip surfaces (MSSs) along principal slip zones in carbonate rocks provide an excellent opportunity to investigate the deformation processes that occur in relation to slip events in the seismogenic zone. The defining feature of MSSs is the high degree of visible light reflectance resulting from a low surface roughness. MSSs can form at seismic slip velocities and, therefore, may indicate paleo-seismicity (Siman-Tov et al., 2013; Smith et al., 2013; Fondriest et al., 2013; Kirkpatrick et al., 2013; Spagnuolo et al., 2015). However, MSS have also been developed at sub-seismic slip conditions in deformation experiments (Verberne et al., 2014) and, thus, they are not necessarily diagnostic of seismic slip.

Studies suggest that the low surface roughness of MMS may be the result of nano-sized grains (< 1 μm), which constitute the uppermost layers of principal slip surfaces (Fondriest et al., 2013; Siman-Tov et al., 2013; Collettini et al., 2014). Therefore, the production of a nanogranular slip-surface coating has been associated with the development of natural MSSs during high-
magnitude seismic events. In addition, nanogranular coatings are also produced in deformation experiments under seismic conditions (Green et al., 2015; Spagnuolo et al., 2015), supporting this hypothesis. Current research suggests different formation mechanisms for MSSs related to nanograins: (1) Formation of long beams by plasticity and subsequent brittle fracturing of the beams (Siman-Tov et al., 2013), (2) localised dynamic recrystallisation and static recrystallisation (Smith et al., 2013), (3) dislocation cell formation (Verberne et al., 2013) and (4) sharp rheological boundaries separating high-, and low-strained regions (Pozzi et al., 2018).

One of the main mechanisms of fault gouge formation is grain-size reduction by comminution. The grain-size dependence of the modified Hall-Petch relation proposed by (Sammis and Ben-Zion, 2008) illustrates that smaller grain sizes require higher differential stresses to fracture. Therefore, local stress concentrations during coseismic events may be favourable sites for generation of nanograins by fracturing micron to millimetre-sized grains. Ultimately, these nanograins may not only be evidence for paleo-seismic events but may in addition control the frictional behaviour of the fault (Han et al., 2007; Han et al., 2010; Di Toro et al., 2011).

Less considered effects of fault rock deformation in carbonates are chemo-mechanical transitions, e.g., decarbonation. Products of decarbonation processes are reported from experiments (Verberne et al., 2014; Spagnuolo et al., 2015; Delle Piane et al., 2017) and from natural faults (Vigano et al., 2011; Collettini et al., 2013). The experimental studies report the presence of amorphous carbon as a deformation product. Some natural faults also exhibit amorphous glass coatings around euhedral calcite crystals in potential carbonate pseudotachylites (Vigano et al., 2011). Furthermore, deformation experiments on silicates (Di Toro et al., 2004; Pec et al., 2012; Hirose et al., 2012; Toy et al., 2015) and on clay-bearing materials (Goldberg et al., 2016) produce non-crystalline or partly-amorphous silicate materials (Pec et al., 2012).
We investigated the surface and the first tens of microns of two natural carbonate fault-mirror structures to test the hypothesis of a nanograin surface coating. Our high-resolution electron microscopy and Raman spectroscopy analyses demonstrate that a thin coating of decarbonation products is able to produce a MMS without the need of a nanogranular surface coating. In addition, we suggest that the amorphous products may facilitate and lubricate seismic slip (De Paola et al., 2011; Di Toro et al., 2011) or possibly re-strengthen the fault by recrystallisation of the decomposition products during the interseismic period.

2. Geological Setting

2.1 Geology of the Arkitsa fault zone

The 700 m long Arkitsa fault (Fig. 1A) belongs to the Kamena Vourla fault system; a northward-dipping, ESE-WNW striking, active normal fault zone of approximately 50 km length along the southern coast of the Gulf of Evia. The left-stepping fault system consists of the Kamena Vourla, the Agios Konstantinos and the Arkitsa fault segments (Roberts and Jackson, 1991; Ganas et al., 1998). The study area (38°43'56.17"N, 23° 0'27.41"E) is situated within the Pelagonian zone of Central Greece. The Arkitsa fault planes juxtapose Late Triassic to Middle/Late Jurassic platform carbonates in the footwall with lower Pliocene-Pleistocene to Quaternary sediments in the hanging wall (Kokkalas et al., 2007). The Arkitsa fault scarp is an anthropogenically formed fault exposure first described by Jackson and McKenzie (1999). Quarrying for two decades removed a major part of the hanging-wall colluvium to reveal three large, smooth fault planes of up to 65 m height (Kokkalas et al., 2007).

Historical seismic records from the Gulf of Evia area record about 13 rupture events from 426 BC until the last major event in 1894 (Ganas et al., 1998; Ganas et al., 2006), with a magnitude Mₗ 6.9 earthquake hosted within the Atalanti fault zone (Fig. 1A) (Ambraseys and Jackson, 1990). This number of events contrasts with the approximately 50 slip increments
identified by Jackson and McKenzie (1999), where the authors calculated the slip increments from fault plane height and average earthquake slip distances. Evidence for Holocene seismic activity along the Arkitsa fault planes is recorded by the approximately 1 m unweathered fault scarp existing prior to quarrying (Jones et al., 2009).

**Figure 1:** Tectonic maps of the study areas. Insets show the geographical location within central Greece. **A:** Location of the studied exposure of the Arkitsa fault zone (modified after Jones et al., 2009; earthquake data: NOAA). **B:** Outcrop location of the Schinos fault segment north of Corinth (modified after Kaplanis et al., 2013 and Collier et al., 1998).

### 2.2 Geology of the Schinos fault zone

The Schinos fault zone is located north-east of Corinth within the Gerania mountains with the studied exposure (38° 2'14.40"N, 23° 0'22.33"E) belonging to the Gerania unit of the Internal Hellenides (Kaplanis et al., 2013). The stratigraphic succession of the Gerania unit includes from bottom to top: Permian sedimentary and volcanic rocks, Upper Triassic neritic limestones and dolomites, Lower Jurassic limestones, an Upper Jurassic succession of rift-related sediments overlain by either Ammonitico Rosso (Bathonian or Oxfordian age) or radiolarian sediments, and ophiolite-derived turbidites of the Beotian flysch (Kaplanis et al., 2013).

Historic seismic data record an earthquake sequence in the Corinth region with three main shocks: 24th February 1981, Ms 6.7; 25th February 1981, Ms 6.4 and 4th March 1981, Ms 6.4 (Collier et al., 1998). Based on the epicentral location and a focal mechanism mismatch, Collier et al. (1998) interpreted that the 24th February event started with displacement on an offshore
fault and continued to rupture the Pisia fault, producing a surface break. In addition, Collier et al. (1998) propose the same succession of events for the 25\textsuperscript{th} February event, which caused a displacement on the Schinos fault, probably forming the most recent event on the fault plane that we study here.

3. Methods

We analysed drill-core samples from the Arkitsa and Schinos fault exposures using a range of micro-analytical techniques. In total, 40 drill cores, 2.54 cm in diameter, were collected from the Arkitsa fault zone and 18 were collected from the Schinos fault zone. The maximum retrievable drill-core lengths did not exceed 10 cm (Arkitsa) and 5 cm (Schinos) as the porosity of the fault rock increases drastically and the material loses its cohesion with distance from the fault surfaces. Seven samples from the Arkitsa fault zone and four samples from the Schinos fault zone were chosen for further investigation based on the best-preserved mirror surfaces. Sample cores were sputter coated with an 8-nm thick layer of platinum/palladium for charge deduction.

From both faults, two representative samples were selected for electron-transparent foil preparation using a FEI Helios Nanolab G3 Dualbeam focused ion beam scanning electron microscope (FIB-SEM). Prior to ion beam deposition of the main platinum strip, a 200-nm layer of platinum was deposited using the electron beam (2kV, 0.4 nA) to prevent surface amorphisation by the ion beam as a preparation artefact. Transmission electron microscope (TEM) investigations of the foils were carried out with a FEI Talos F200X. TEM images were acquired with 200 kV acceleration voltage and 5 nA or 10 nA beam current depending on final sample thickness. The FEI Talos SuperEDX detector was used for chemical element mapping via energy dispersive X-rays (EDX) in scanning transmission electron microscope (STEM) mode. EDX maps were quantified post-acquisition with Bruker Esprit 1.9 software using the Cliff-Lorimer approximation and ImageJ.
Electron energy loss spectroscopy (EELS) data were acquired with a Zeiss Libra 200FE in TEM mode at 200 kV with an Omega in-column energy filter. The energy resolution of EELS analyses was 0.7 eV, measured at the full width half maximum (FWHM) of the zero-loss peak. Energy loss spectra were obtained at 250,000x magnification with a 100 μm filter-entrance aperture giving an effective aperture of about 40 nm on the sample. The convergence angle of EELS was about 0.1 milliradian (mrad), and the acceptance angle was defined by the 60 μm diameter of the objective aperture giving a collection angle of 11.6 mrad. Measurement times of EEL spectra were set to 1–5 s, with 5–10 frames/spectra, on a slow-scan CCD (model UltraScan 4000), with binning of 4x4 pixels (giving 1024 pixels), and an energy spread of 0.08 eV/pixel, resulting in an energy range of 82 eV in the acquired spectra. Spectra were deconvolved with the zero-loss peak to remove the effect of plural scattering and background-subtracted assuming a power law function with Gatan’s Digital Micrograph software.

Raman spectroscopy was carried out with a WiTec ALPHA300 R confocal microscope to identify and characterize possible reaction or decomposition products on the fault surfaces. We used a laser with a wavelength of 532 nm and a spectral grating of 600 grooves/cm. Data acquisition, data post-processing and peak fitting were performed with WiTec ProjectFour 4.1 software and Fityk 0.98 (Wojdyr, 2010) peak fitting software. In accordance with (Ferrari and Robertson, 2000) we determined the intensity ratios of the D to G peak, \( \frac{I(D)}{I(G)} \), to obtain information about the degree of crystallinity within carbon phases. We used the approach by Ferrari and Robertson (2000) based on first principle considerations to determine the peak height ratios because the information about the less disordered aromatic rings and the clustering of the \( \text{sp}^2 \) phase is contained in the intensity maximum of the D peak and not in the width. Calculation
of reaction enthalpy values were executed with SUPCRT92 (Johnson et al., 1992) at standard
conditions (1 bar, 298 K).

4. Results
4.1 Field results
The exposed fault surface of the Arkitsa fault (Fig. 2A) exhibits a range of slip-related
structures. On the outcrop scale, the fault exposure steps over, suggesting the presence of several
slip planes inside the fault damage zone. The fault surface contains pronounced slip grooves
parallel to the direction of oblique slip. Fractures with a spacing on the order of one metre are
oriented approximately perpendicular to the grooves (Fig. 2A). Parts of the slip plane are covered
with residual reddish-brown hanging-wall breccia, incorporating fragments of dark host-rock
carbonate up to several decimetres in size (Fig. 2A). The most prominent feature is the low
roughness of the fault surface, which enables the reflection of sunlight (Fig. 2B). The fault rock is
a matrix-supported greyish, cohesive material, with dark carbonate clasts up to several
centimetres in size (Fig. 2C). Slip-parallel alignment of clasts on the fault surfaces can be traced
over several meters (Fig. 2C).

The Schinos fault surface (Fig. 2D) exhibits a range of structures formed during slip. We
observe stepovers, indicating the presence of several fault planes inside the fault damage zone.
The fault surface is extremely smooth with areas not only reflecting light but also mirroring the
surrounding vegetation (Fig. 2E). Fragments of grey carbonate host rock are incorporated into the
red-orange footwall fault rock. The clasts are strongly aligned in the slip direction and usually do
not exceed 1–2 cm in size (Fig. 2F). The fault surface contains pronounced slip grooves and a
wavy surface morphology (Fig. 2D). The mirror surfaces of both faults are hosted inside a zone
of high competence with average thicknesses of 10 cm and 5 cm for Arkitsa and Schinos,
respectively.

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Figure 2: Representative field structures from Arkitsa (A – C) and Schinos (D – F) fault exposures. A: Overview of Arkitsa fault exposure showing one of the three main slip planes with reddish-brown hanging-wall breccia cover on part of the slip surface. Black arrows indicate fractures perpendicular to slip direction. Person for scale, bottom right corner. B: Fault surface reflecting sunlight. C: Alignment of host rock fragments parallel to slip direction. Box approx. 5 x 4 cm. D: Schinos fault surface exposure shows grey weathering of the slip plane. The slip plane is curved with the salient pointing out of the figure. Fault scarp to the north is now covered by alluvium. E: Image demonstrating fault surface reflectivity. Reflection of vegetation along the top edge and above the one-euro coin. F: Grey, weathered fault plane showing a strong alignment of light grey host-rock clasts parallel to slip direction (one-euro coin for scale, bottom edge).

4.2 Slip surface microstructures

SEM analyses of the Arkitsa and Schinos slip surfaces reveal a low surface roughness on the microscopic level. The surfaces are well polished and, in some places, preserve parts of the
hanging wall (Fig. 3A and F). Microgrooves are aligned parallel to the slip direction (Fig. 3B) and holes with a diameter of 6–8 μm pierce the slip surfaces (Fig. 3A and D). Localized erosion of the Arkitsa slip surface exposes the underlying fault rock, with a grain size of 2–5 μm (Fig. 3A, B and C). The first 10–20 μm of the fault rock below the Arkitsa slip surface consist of a fine-grained deformation product, which is situated on top of a comparatively coarse-grained, less deformed material (Fig. 3B and C). The contact between the fine- and coarse-grained domains is established by a coating phase (Fig. 3B and C).
Figure 3: Representative slip surface microstructures of the Arkitsa (A-C) and Schinos (D-F) exposures. A: Secondary electron overview image of the Arkitsa slip surface. Smooth slip patches with microgrooves parallel to slip direction. Uniform holes (white arrows) penetrate the slip surface. Residue of hanging wall on top of the slip surface patch (black arrows and inside the white circle) with TEM sample location. B: Close-up of slip surface from A. Fine-grained slip surface material situated on top of coarse fault rock grains. Microscope stage tilted to 52°. C: Close-up of section from B. Roundish grains are covered by coating phase (white arrows). D: Secondary electron image of Schinos slip surface with microgrooves parallel to slip direction. Uniform holes penetrate the principal slip surface (black arrows). Inset: Backscatter electron image illustrating triple junction grain boundary contacts. Scale bar = 50 μm. E: Backscatter electron image top view onto the slip surface. Cracks in idiomorphic calcite crystals are visible a few micrometres into the crystal (black-white arrows). Coating phase with low backscatter contrast covers parts of the slip surface and reduces surface roughness (white arrows) F: Backscatter electron image of the contact between footwall and hanging-wall. Black arrows indicate locations of a coating phase with lower backscatter contrast.

In contrast, the Schinos fault surface consists of large, truncated grains with boundaries that meet in triple junctions (Fig. 3D and inset). A fragmented layer lies on top of undeformed calcite crystals (Fig. 3E) and the damage extends about 10–20 μm into the fault rock. Here also a coating phase is present on the slip plane (Fig. 3E and F). On both faults, Arkitsa and Schinos, the coating phase has a low contrast in backscattered-electron images (Fig 4A and B). The phase also coats grain edges and reduces the slip surface roughness (Fig. 3C, E and F). In addition, the coating phase infiltrates interstitial regions of the hanging-wall breccia and holds residual pieces of hanging wall in place (Fig. 3F).
Figure 4: Backscatter electron images with locations of Raman spectroscopy measurements for Arkitsa (A and C) and Schinos (B and D) slip surfaces. A and B: Backscatter electron images of the fault surfaces. A phase with low backscatter contrast (darker grey) is visible in B. C and D: Raman spectra showing the D and G peak position of a disordered carbon phase.

Figure 4 presents Raman spectra from the coating phase. Two broad peaks between 1200–1700 cm\(^{-1}\) indicate the presence of a disordered carbon phase, whilst a peak at 1086 cm\(^{-1}\) demonstrates the presence of crystalline calcite. The fitted spectrum from the Arkitsa fault plane gives a \(\frac{I(D)}{I(G)}\) ratio of 0.543 at a D-peak position of 1339 cm\(^{-1}\) and a G-peak position of 1584 cm\(^{-1}\).

The fitted spectrum from the Schinos fault plane gives a \(\frac{I(D)}{I(G)}\) ratio of 0.789 at a D-peak position of 1344 cm\(^{-1}\) and a G peak position of 1587 cm\(^{-1}\).

4.3 Nanostructures

TEM analyses on FIB-SEM foils of both fault-surface exposures reveal a thin surface coating. The uppermost layer of both fault-rock exposures is defined by a principal slip surface with a thin, non-crystalline coating between hanging wall and footwall (Fig. 5A and B). The coating has a homogenous diffraction contrast, shows no lattice fringes (Fig. 6A to D, 8A and F) and EDX analyses indicate the presence of carbon (C), iron (Fe), silicon (Si) and aluminium (Al) (Fig. 7, Si content of Arkitsa ~ 40 area % estimated with ImageJ). The coating connects hanging-wall breccia with the footwall fault surface (Fig. 6B and D) but also extends into the hanging-wall breccia (Fig. 6A and C). EELS measurements of the coating phase are given in the inset of Fig. 6D. The onset of the \(\pi^*\) peak represents the measured carbon K-edge at 284 eV which is followed by a ‘whaleback-shaped’ \(\sigma^*\) region. In addition, the measurement of the Schinos sample exhibits an additional peak at 286.5 eV possibly indicating C-O bonding.
Figure 5: Overview of TEM cross sections into the principal slip surfaces of Arkitsa and Schinos, including preserved hanging-wall residue. **A:** Bright-field TEM image overview of representative Arkitsa nanostructures. The hanging-wall breccia consists of small, elongated calcite fragments held in place by non-crystalline carbon. The nanogranular layer is part of the hanging-wall breccia and lies on top of deformed calcite crystals. Fractures (dashed lines) bend down from the principal slip surface into the deformed calcite crystals. Fractures and holes form porosity inside the mirror slip volume and are filled either with non-crystalline carbon and/or nano-clay. **B:** Bright-field TEM image with representative Schinos nanostructural hanging-wall to footwall relation. The hanging wall consists of small grains of several hundred nanometres. Individual grains are surrounded by non-crystalline carbon. The footwall hosts a nanograin filling, the grain size of which increases abruptly with distance from the slip surface to grains of several micrometres in size. Larger grains contain twins. hw = hanging wall, fw = footwall.
We observe a direct relation between the spatial distribution of amorphous material and calcite nanograins. For the Arkitsa fault exposure, a nanogranular layer is present with rounded, elliptical calcite nanograins of ~50 nm size between the footwall and the hanging-wall breccia (Fig. 5A, 6A, B and 7). Selected area electron diffraction (SAED) patterns of the Arkitsa nanogranular layer in Figure 5A demonstrate the polycrystalline nature of the material with clustered diffraction spots forming two arcs in an angle of about 75° (inset Fig. 6A). Figure 8A reveals that the thickness of the coating varies along the slip surface and can locally incorporate calcite grains from the underlying nanostructure. We find that calcite nanograins are not restricted to the principal slip surface but also occur away from the displacement zone inside the fault rock (Fig. 8B). The nanograins do not contain any dislocations (Fig. 6B and 8B) and are not always in grain-to-grain contact in the plane of the section. Calcite nanograins of the Schinos exposure are mostly located between large grains along the principal slip surface (Fig. 8C and D). The gaps between large grains are filled with an amorphous material that increases the overall smoothness of the fault surface. Inside the amorphous material, most nanograins have elongated shapes, with no dislocations structure and nano-clay minerals often wrap around the grains (Fig. 8D).
Figure 6: TEM cross sections illustrating the relationship between hanging-wall (hw) breccia and footwall (fw) contact for Arkitsa (A – B) and Schinos (C – D). A: TEM bright field close-up from Fig. 5A. Non-crystalline carbon establishes the contact between hanging wall and footwall. The footwall consists of large, stressed calcite crystals with bent fractures. Nanogranular layer with non-crystalline carbon and calcite nanograins. White circle indicates SAED location; black circle depicts EELS measurement. Inset: SAED showing two sets of crystal orientation. ss = slip surface. B: TEM bright field close-up from A. Non-crystalline carbon connects hanging wall and footwall. The nanogranular layer produces Moiré fringes indicative of overlapping crystal lattices. Nanograins are not always in grain-to-grain contact. C: TEM bright field close-up image from Fig. 5B. The grain size of the hanging wall is 200 nm and larger. The grain size of the footwall directly at slip surface is about 100 nm. D: TEM bright field close-up image from C. The hanging wall and footwall are connected via non-crystalline carbon. Black circle depicts EELS measurement. Inset shows EELS spectra from black circles in A and D.
Further TEM investigation of the slip surfaces reveals a variety of slip-derived deformation nanostructures. Neither fault exposure exhibits a gradual decrease in grain size towards the slip surface but show an abrupt change, instead. Adjacent to the principal slip surface, we observe twinned calcite crystals about 2–5 μm in size (Fig. 5A and B). Dislocations are homogeneously distributed in the larger crystals from the Arkitsa exposure (Fig. 5A). For the Schinos case, the dislocation distribution appears to be less ordered, with dislocations...
concentrated towards grain contacts in the slip direction (Fig. 8C). Bent cleavage fractures dip
down from the Arkitsa slip surface and exhibit minor displacements together with slight increases
in dislocation density along the fracture planes (Fig. 5A and 6A). Along the slip surface, the
nanostructure of the Arkitsa exposure can vary in intensity and develop a foam nanostructure
(Fig. 8A). The layer thickness of the foam nanostructure is about 1 μm and the grain size ranges
from 100–200 nm with the grain boundaries exhibiting triple junctions at ~120° angles (Fig. 8A).
This nanostructure is not observed in the samples analysed from the Schinos exposure, which
instead contain dislocation arrays (Fig. 8C and E). In general, the foam nanostructure at Arkitsa
appears to be less developed adjacent to the slip surface, especially where the fracture density is
highest.
Figure 8: Nanostructures of fault surface coating and nanograin abundance not associated with the principal slip surface. A: Bright-field TEM image from the Arkitsa exposure. Amorphous slip surface coating with calcite nanograins and crystal fragments from underlying foam nanostructure (black arrow). A sharp contact separates the underlying nanostructure from the slip surface coating. Nanostructure with former fracture planes (dashed lines), which originate from the principal slip surface, and overprint the existing nanostructure. B: Calcite nanograins away from the principal slip surface hosted inside non-crystalline carbon. Tip direction of black-white arrows indicate lattice fringe orientation of new grains. C: Bright-field STEM image overview. Large grains (5 μm) with a heterogeneous dislocation density distribution. D: Calcite nanograins in non-crystalline carbon as a filling between two grain boundaries directly at the principal slip surface. Clay minerals wrap around the nanograins (black arrows). E: Bright-field STEM close-up from C showing the dislocation structure directly at the principal slip surface. Black arrows point to dislocation-pinned Frank-Read sources frozen during dislocation bow-out. F: Bright-
field TEM image from the Schinos exposure. Amorphous slip surface coating mixed with nano-clay on top of calcite crystal. cc = calcite.

5 Discussion
5.1 Processes revealed by nano/microstructures
5.1.1 Decarbonation and carbon reduction

Nanostructural investigation of the Arkitsa and Schinos fault exposures reveal the importance of calcite decarbonation products for the formation of a smooth fault surface. TEM images reveal a coating of an amorphous solid that we termed “coating phase” or “amorphous material” above. We suggest that the amorphous solid formed following calcite decarbonation, as a result of shear heating at asperity contacts or the 'severe' introduction of dislocations during fault slip. The dislocation production takes place via (re-)activation of Frank-Read sources, which can multiply dislocations and emit dislocation loops into the crystal after dislocation bow-out. The Schinos fault exposure preserves an example of two activated Frank-Read sources close to the principal slip surface (Fig. 8E). In our natural samples, holes piercing the principal slip surface (Fig. 3A and D) indicate CO₂ degassing as a product of the decarbonation reaction

$$\text{CaCO}_3(s) \rightleftharpoons \text{CO}_2(g) + \text{CaO}(s) \quad (\Delta H^{0}_{298 K} = +178 \text{ kJ mol}^{-1}) \quad (1)$$

reported to occur due to thermal decomposition of calcite starting at 600 °C (Rodriguez-Navarro et al., 2009). However, experiments by Martinelli and Plescia (2004) recorded CO₂ emission during ball milling experiments of calcite without increasing the macroscopic temperature. In addition to the CO₂ emission, the authors inferred the presence of amorphous lime (CaO) in their milling product as a result of a process termed ‘mechanical liming’. Subsequent heating of the milling product resulted in a decrease of amorphous lime and the formation of calcite. The experiments of Martinelli and Plescia (2004) demonstrate that reaction (1) can also take place just by deforming calcite without reaching macroscopic temperatures of > 600 °C. Based on these experiments, we suggest that the lime produced during coseismic decarbonation may be partly
amorphous. The experiments also suggest that decarbonation can occur at macroscopic
347 temperatures lower than 600 °C, although the temperature at asperity contacts could be higher.
349 We suggest that 'mechanical liming' is important for wet deformation conditions where fluids are
350 present and will buffer the temperature increase.

The acquired electron energy loss and Raman spectra indicate the presence of a carbon
351 phase. Hence, the presence of this phase implies the occurrence of carbon-reducing processes
352 because reaction (1) does not produce elemental carbon. Precipitation of carbon in deformation
353 experiments of Oohashi et al. (2014) under a hydrogen atmosphere suggests that the reaction
354
\[ \text{CO}_2(g) + 2 \text{H}_2(g) \rightarrow \text{C}(s) + 2 \text{H}_2\text{O}(g) \quad (\Delta \text{H}^0_{298K} = -90 \text{kJ mol}^{-1}) \] (2)
355
356 can reduce CO\(_2\) to form carbon. Similarly, Spagnuolo et al. (2015) detected H\(_2(g)\) and CO\(_2(g)\)
357 degassing during their experiments. Although reaction (2) is expressed as a chemical vapour
358 deposition reaction some of the hydrogen produced can go into solution to reduce dissolved CO\(_2\).
359 This process is known to occur in hydrothermal systems where dissolved hydrogen can interact
360 with CO\(_2(aq)\) to form reduced carbon phases (e.g., (Milesi et al., 2015)). Under wet conditions, the
361 CO\(_2\) from reaction (1) goes into solution starting the hydrolysis
362
\[ \text{CO}_2(g) \rightleftharpoons \text{CO}_2(aq) + \text{H}_2\text{O} \rightleftharpoons \text{H}_2\text{CO}_3 \]
363
\[ \rightleftharpoons \text{HCO}_3^- + \text{H}^+ \]
364
\[ \rightleftharpoons \text{CO}_3^{2-} + 2 \text{H}^+ \]
365 to give carbonic acid, hydrogen carbonate, carbonate ions, and protons (Ruiz-Agudo et al., 2013).
366 The mole fraction solubility (X\(_1\)) of CO\(_2\) in water at room temperature is, with X\(_1\) = 6.15 \times 10\(^{-4}\),
367 one order of magnitude larger than H\(_2\) with X\(_1\) = 1.411 \times 10\(^{-5}\) (Gevantman, 2000). It is, therefore,
368 more likely that a larger amount of CO\(_2\) will dissolve in water and form carbonic acid, being the
369 more probable proton donor, for reducing carbon from the residual CO\(_2\). As an alternative, CaO
surfaces can act as potential catalytic sites and promote the reaction with H₂O at kinks, corners and/or steps. These surfaces can split H₂O leading to one hydroxyl ion (OH⁻) and one proton (H⁺) which in turn may also be able to reduce carbon from CO₂ (Kudłacz and Rodriguez-Navarro, 2014). The main requirement for this reaction to occur is the presence of crystalline CaO. This would either require prior crystallisation of the suspected amorphous lime from reaction (1) or the lime resulting from reaction (1) was already crystalline. Calcium ions (Ca²⁺) decrease the pH of the water facilitating CO₂ solubility. Earth-alkali metal reactions with water are commonly known to produce hydrogen by reaction of the metal with water to form H₂ and the corresponding metal hydroxide. As the reduction of CO₂ by hydrogen does not require the participation of an additional reducing agent, it is likely to occur also under closed-system experimental conditions. With a variety of possible production reactions, it remains challenging to identify the main reaction responsible for the source of hydrogen for CO₂ reduction. However, a combination of several processes is likely, with the simplest one being the hydrolysis of CO₂ in H₂O. The CaO or Ca²⁺ interaction with water can increase the pH up to 12.4, as shown in experiments by Ruiz-Agudo et al. (2013), which increases the solubility of CO₂ in water and would facilitate reaction (3). In general, the result would be the precipitation of carbon from the fluid, which would not be restricted to the principal slip surface.

5.1.2 Carbon-bond sp² hybridisation

The obtained \( \frac{I(D)}{I(G)} \) ratios from Raman spectroscopy indicate that the slip surfaces are coated with nanocrystalline graphite according to the interpretation of the Raman modes by Ferrari and Robertson (2000). With the obtained \( \frac{I(D)}{I(G)} \) ratios we can calculate the size of the nanocrystallites according to the equation from (Pimenta et al., 2007)
where \( L_a \) is the cluster/crystallite size in nm, \( E_{\text{laser}} \) is the energy (wavelength) of the laser with 2.33 eV (532 nm), and \( \frac{I(D)}{I(G)} \) the intensity ratio of D and G peaks. Equation (4) is based on results from Pimenta et al. (2007), which revealed an inversely proportional relationship between the fourth power of \( E_{\text{laser}} \) and the \( \frac{I(D)}{I(G)} \) ratio. For the Arkitsa exposure we calculate a size of \( L_a = 35 \) nm (\( \frac{I(D)}{I(G)} = 0.543 \)) and for the Schinos exposure \( L_a = 24 \) nm (\( \frac{I(D)}{I(G)} = 0.789 \)). While Pimenta et al. (2007) calculate with \( L_a \) the graphite nanocrystallite size, Ferrari and Robertson (2000) mention \( L_a \) as the cluster diameter of carbon aromatic rings. The terminology used by Ferrari and Robertson (2000) for nanocrystalline graphite and amorphous carbon is based on the starting material experiencing disorder. When disordering graphite, \( \frac{I(D)}{I(G)} \) will increase with increasing disorder and development of a D peak. With amorphous carbon as the starting material, the development of a D peak and the increase in \( \frac{I(D)}{I(G)} \) indicates clustering. Because the D peak arises from aromatic rings, for small \( L_a \) the D-mode strength is proportional to the cluster area or diameter Ferrari and Robertson (2000).

According to our discussion of possible options for carbon reduction in Section 5.1.1, where carbon is most likely produced as the outcome of post-seismic, chemical reduction processes, the initially precipitated carbon may be amorphous. Nathan et al. (1974) reported for glass-like carbon a D-peak position at 1340 cm\(^{-1}\) and a G-peak position at 1590 cm\(^{-1}\), while Wang et al. (1990) reported a D-peak position at 1347 cm\(^{-1}\) and a G-peak position at 1588 cm\(^{-1}\). The band positions from Wang et al. (1990) were obtained with a 515 nm laser on glass-like carbon, which was heat treated at 2000 °C. Wang et al. (1990) also noted that the D-peak position is
sensitive to the wavelength of the laser used, exhibiting a band-position shift towards lower wavenumbers with increasing laser wavelength. The reported values from Nathan et al. (1974) and Wang et al. (1990) are in agreement with our measured band positions for Arkitsa (D peak: 1339 cm\(^{-1}\), G peak: 1584 cm\(^{-1}\)) and Schinos (D peak: 1344 cm\(^{-1}\), G peak: 1587 cm\(^{-1}\)), suggesting that the slip surface coating resembles glass-like carbon. Therefore, instead of the development of nanocrystalline graphite, our results are consistent with the presence of a partly-ordered carbon exhibiting an electron-bond environment akin to glass-like carbon. This may be evidence for the beginning of a clustering process of the aromatic carbon rings (sp\(^2\)-clustering), which is likely to have occurred during post-seismic annealing.

High-resolution TEM imaging coupled with electron diffraction suggests that the surface coating is non-crystalline. This interpretation is supported by our EELS analysis which results in spectra resembling amorphous carbon (Fig. 9). However, although both have a \(\pi^*\) peak, the phase identified here has a more pronounced \(\sigma^*\) region indicating a stronger graphitic order, yet lacking full long-range order as observed in crystalline graphite (Rosenberg et al., 1986). Our spectra resemble those of Stroud et al. (2011), suggested to be indicative of glass-like carbon with sp\(^2\) hybridisation. Although the EEL spectra resemble glass-like carbon, we will refer to the carbon phase from the fault exposures as partly-hybridised amorphous carbon (PHAC). Impurities of Al, Fe, and Si (Fig. 7) in the PHAC suggest either the concomitant decomposition of silicate phases (e.g., clays) during slip or the influence of hydrothermal fluids percolating along the fault zone, precipitating clay. We suggest that the presence of PHAC demonstrates that amorphous carbon is precipitated during or immediately after slip but undergoes ordering (sp\(^2\)-hybridisation) via post-seismic annealing, as demonstrated by Raman spectroscopy and EELS.
Figure 9: Comparison of electron energy loss spectra. Calcite decomposition products presented here have a small intensity difference of $\pi^*$ to $\sigma^*$ peak. The EEL spectrum from vitreous (glass-like) carbon (Stroud et al., 2011) exhibits a similar $\sigma^*$-peak shape compared to the acquired spectra of this study. Reference spectrum of graphite has three weak $\sigma^*$ peak features which are absent in all other carbon spectra.

5.1.3 Nanograin nucleation

Our nanostructural investigations reveal a close spatial relationship between PHAC and newly formed calcite nanograins (e.g., Fig. 6A). Nucleation of nanograins inside the decomposition product implies a similar chemistry of reactant and product. As discussed in Section 5.1.1, reaction (1) also produces lime, which is highly reactive and can form portlandite (Ca(OH$_2$)) under hydrous conditions via the hydration reaction

$$\text{CaO}_{(s)} + \text{H}_2\text{O} \rightarrow \text{Ca(OH)}_{2(s)} \quad (\Delta H^0_{298K} = -65 \text{ kJ mol}^{-1}) \quad (4).$$

In the presence of CO$_2$, portlandite back-reacts to calcite and water by the carbonation reaction

$$\text{Ca(OH)}_2 + \text{CO}_2 \rightleftharpoons \text{CaCO}_3 + \text{H}_2\text{O} \quad (\Delta H^0_{298K} = -93 \text{ kJ mol}^{-1}) \quad (5).$$
Kudlacz and Rodriguez-Navarro (2014) observed a nucleation-related crystallographic preferred orientation in two sets of portlandite crystals. The angle between the \( \{11-20\} \) portlandite planes was 75°, implying that portlandite developed with a topotactic relationship of the \( \{11-20\} \) portlandite planes parallel to the \( \{110\} \) \( \text{CaO} \) planes of lime. The SAED inset of Fig. 6A indicates two preferred orientations with an angle of about 75° (centre to centre) between the \( \{11-20\} \) planes of calcite in our natural samples, supporting our described back-reaction steps from lime, through portlandite, to calcite. In addition, we also observe dislocation-free nanograins away from the slip surface (Fig. 8B). Hence, we suggest that calcite nanograins are not the outcome of comminution or plastic grain-size reduction during slip but originate from pseudomorphic growth of calcite after portlandite and portlandite after lime from decarbonation.

5.2 Fault surface evolution during the seismic cycle

The onset of fault slip triggers a variety of crystal-plastic deformation processes. Cleavage fractures in Figure 3E demonstrate that slip localises within the first 20 μm inside the fault rock. This shows that the transfer of strain from the slip surface is low because the cleavage fractures are concentrated close to the slip surface. Fractures also cross-cut the foam nanostructure and produce a nanofabric where grains are trapped between sheet-like fracture planes (Fig 8A). Shear heating may cause a considerable temperature rise at asperity contacts. The temperature increase or grain fracturing trigger the decarbonation reaction (Eq. 1), which leads to \( \text{CO}_2 \) degassing and the production of lime. It is not fully resolved whether the lime would be amorphous, as reported by Martinelli and Plescia (2004), or crystalline.

During and immediately after fault slip, calcium oxide infiltrates cracks and cavities and reacts with the fluids present to crystalline portlandite (Eq. 4). The resulting portlandite crystals back-react with the released \( \text{CO}_2 \) to form calcite nanograins (Eq. 5). During the pseudomorphic replacement of portlandite by calcite, the calcite nanograins preserve the crystallographic
nucleation relationship from portlandite. Nano-clay precipitates inside the reaction medium either from the fluid or from previously decomposed clay minerals and may contribute to the fault-mirror appearance. Reduction of carbon can occur by reducing CO$_2$ with dissolved hydrogen (Eq. 2) either from the hydrolysis reaction (Eq. 3) or by water splitting with lime as a catalyst. Subsequent sp$^2$-hybridisation of the amorphous carbon results in aromatic ring clustering and hence in a stronger near-field bonding of the carbon. As a result, PHAC is formed.

Graphitisation of carbonaceous materials in active faults may be an important factor for fault reactivation because graphite is characterised by a low friction coefficient of $\mu_{ss} = 0.08$ (Oohashi et al., 2011). Glass-like carbon is a stable, non-graphitising type of carbon which requires an activation energy of +215 kJ mol$^{-1}$ to initiate graphitisation (Saxena and Bragg, 1978; Hokao et al., 2000). This would imply that on natural faults graphite can only be found after a considerable energy input to induce graphitisation of glass-like carbon species. Saxena and Bragg (1978) identified three stages for graphitisation of glass-like carbon: (1) < 1500 °C dehydrogenation, (2) 1500 °C $\leq$ 2300 °C stress relief of the structure and (3) > 2300 °C onset of graphitisation. A high temperature treatment of synthetic glass-like carbon can cause graphitisation suggesting a thermodynamic control of the graphitisation process. Molecular dynamics simulations of sp$^2$ carbon-bond self-assembly illustrate that sp$^2$-clustering from amorphous precursors can commence after 200 picoseconds at $\sim$3200 °C (Powles et al., 2009). This short time span suggests that the sp$^2$-clustering we observe here takes place immediately after carbon reduction. In addition, the simulations suggest that once sp$^2$-bonded stable clusters form, breaking these bonds to further rearrange and increase sp$^2$-ordering may require the need to overcome an additional energy barrier. This could explain why the PHAC observed here never reached a higher degree of graphitisation because the energy input was not enough to break the initial stable carbon clusters. Also, synthetic glass-like carbon displays the often described,
entangled, lath-like structure (e.g., (Saxena and Bragg, 1978)). Although the PHAC observed here lacks the aforementioned nanostructure, it does have the vibrational and electronic properties of glass-like carbon (Fig. 4 C and D, Fig. 9). Because PHAC is formed by chemical precipitation and hybridisation, in contrast to pyrolysis synthesis of glass-like carbon (Powles et al., 2009), we propose that the way the carbon is formed influences the structure of the material while maintaining a similar interatomic bond structure. This suggestion is supported by Powles et al. (2009), who point out that the properties of glass-like carbon depend on the formation conditions, the precursor material, and on the annealing conditions. It is not clear whether the consideration of geological time scales for the annealing of PHAC at lower temperatures may yield a similar result in graphitisation.

5.3 Advances in understanding fault surface evolution

Amorphous carbon has been observed in deformation experiments (Verberne et al., 2014; Spagnuolo et al., 2015; Delle Piane et al., 2017) and its mechanical role has been assessed by Oohashi et al. (2011). The authors measured a friction coefficient for amorphous carbon of $\mu = 0.54$ at the initiation of slip and a low steady-state friction coefficient of $\mu_{ss} = 0.15$ at seismic slip velocities. Di Toro et al. (2011) speculated about fault lubrication by carbonate decomposition products but, for the first time, we can demonstrate the extent of amorphous coatings. Decarbonation is the predominant process for producing a thin fault surface coating and is mainly responsible for the low surface roughness by smoothing out surface corrugations. Because the fault surfaces show little variation in the field, our results suggest that the fault planes of both fault exposures are at present covered with the decarbonation and carbon hybridisation products.

Our EEL and Raman spectroscopy results indicate that PHAC has a similar carbon-bonding environment compared to glass-like carbon. Synthetic glass-like carbon has an entangled lath-like structure consisting of ordered carbon sheets while the natural PHAC has no
nanostructure. The mechanical properties of glass-like carbon, heat treated at 1000 °C, show a
Youngs modulus of $E = 29$ GPa and a shear modulus of $G = 12.5$ GPa (Robertson, 1991). Hokao
et al. (2000) report a friction coefficient for glass-like carbon of $\mu = 0.21$ and $\mu = 0.13$ for
mixtures of glass-like carbon with graphite. However, as the structure of the PHAC is different
from synthetic glass-like carbon we suggest that mechanical properties of the synthetic material
may not fully reflect the mechanical behaviour of the carbon phase reported here. Because little is
known about the deformation of various crystalline and non-crystalline carbon phases further
experimental investigations are warranted.

While previous research focused mainly on grain size reduction mechanisms (Sammis and
Ben-Zion, 2008; Siman-Tov et al., 2013), our results highlight the importance of decarbonation
and back-reactions as part of the overall fault deformation and healing behaviour. In contrast to
(Siman-Tov et al., 2013) we do not identify small grains evolving from fracturing long beams of
calcite. However, our results indicate that, for natural carbonate faults, a nanogranular coating is
not a key feature for producing a fault mirror surface. For example, the Schinos fault exposure
illustrates that MMSs can contain no coating of calcite nanograins (Fig. 7C, E and F). Instead,
MSS are produced by the interplay of grain-size reduction, decarbonation, back-reactions, and
annealing. The present study does not exclude crystal-plastic deformation as an important grain-
size reduction mechanism in fault gouges but demonstrates that it is not the main mechanism to
form a fault mirror surface. The formation of an amorphous phase with low viscosity during
deformation could explain the low friction, stable, near steady state behaviour after the onset of
weakening in Han et al. (2007) as well as the increase in friction coefficient (Spagnuolo et al.,
2015) via re-carbonation and carbon hybridisation after displacement stopped. The proposed
chemical reactions demonstrate that a succession of healing reactions take place after fault slip
ceases.
6 Conclusions

We conclude that decarbonation of calcite and the subsequent reaction of the decarbonation products produces fault mirror surfaces. The decarbonation process itself may be a major coseismic fault weakening factor and fault slip is facilitated on a decarbonation-product glide film. Frictional behaviour during slip may be dictated by lubrication of low viscosity (amorphous) calcium oxide and (amorphous) carbon. Occasional precipitation of clay inside the decarbonation products. Post-seismic hybridisation produces partly-hybridised amorphous carbon and connects footwall with hanging wall. Ultimately, post-seismic carbonation of portlandite produces new nanograin calcite crystals by pseudomorphic replacement and illustrate the fault healing.

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