# Use of Dyes to Detect Fingermarks

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Use of stains to detect fingermarks

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Abstract

Detection of fingermarks at a crime scene or on related items is of prime interest for forensic investigators, mainly for identification purposes. Most of the fingermarks are invisible to the naked eye, however, and application of detection techniques is required to establish visual contrast between the secretion residue and the underlying substrate. We give here a review of the field related to the concept of using stains to detect fingermarks. A distinction has been made between the physically driven classical detection techniques, the chemically driven ones, and those based on nanostructured materials, an emerging field in forensic science.

Key words: amino acid, blood, detection method, dye, fingermark, fingerprint, forensic, science, identification, lipid, luminescence, nanoparticle, powder, trace evidence
In a criminal investigation, one role of the forensic scientist is to collect information that permits the identification of people involved in the events. Fingerprints constitute one of the most important items of physical evidence for identification. A fingerprint is defined as a unique pattern present on the fingertips of each individual as well as on the palms and the soles. A fingerprint is characterized by invariability from birth to death, excluding scars and irreversible skin damage that could occur during one's life. In this context, we chose to make a distinction between fingerprints and fingermarks. Fingerprints are physically present on the individual's body and can be inked on a ten-print card to be used for comparison in an identification process. Fingermarks are defined as the secretion residue that is left by a person when a surface is touched with unprotected hands. The origin of this residue is in the pores along the skin surface through which human sweat is excreted. The chemical composition of sweat varies according to the nature of the secreting gland and its location on the body (Ramotowski 2001). In addition, contaminants such as cosmetics or other exogenous components can be present on the fingertips due to contact of bare hands with one's face or fatty food. As a consequence, the composition of the residue that is left on a substrate generally is a complex mixture of organic and inorganic materials dispersed in an aqueous matrix, e.g., amino acids, fatty acids, glycerides, inorganic ions, proteins and esters (Ramotowski 2001). Most importantly, such residue generally is not visible to the naked eye. In such situations the fingermarks are said to be "latent" and the application of detection techniques is required to permit their visualization (Champod et al. 2004). This constitutes a major and continuing challenge for forensic scientists.

We address here the different techniques and reagents used to visualize latent (invisible) fingermarks. The key principle of fingerprint detection is simple: to generate as great an optical contrast as possible between the ridges constituting the fingerprint pattern and the underlying substrate without disturbing the pattern formed by the trace. This is necessary
to ensure efficient visualization of the latent fingermarks, their ridges and their characteristics so they can be used for identification. For white or transparent substrates (e.g., printer paper or polypropylene plastic sheets), good contrast can be obtained readily by staining the residue using dark colored dyes. When dealing with colored, patterned or complex substrates, e.g., magazines, commercial packages and banknotes, however, dyes are not appropriate, and the use of a luminescent stain is necessary to avoid confusion due to the background pattern. The use of luminescence to detect fingermarks also is generally preferred for sensitivity reasons. Actual fingermarks sometimes are composed of extremely small amounts of secretions due to a lack of secretion deposited or to further alterations caused by time and environmental conditions. This makes them difficult to detect when stained with classic chromophores. The luminescence mode, however, increases the sensitivity of detection so that faint fingermarks can be recorded and processed for identification. This also explains why current research projects in the field are directed primarily toward development of new and efficient luminescent detection techniques.

To obtain good contrast, it is necessary for a stain to interact specifically with the secretion residue and not with the underlying substrate (or reciprocally, but this is less often the case). Because the composition of fingermarks varies according to various parameters that forensic scientists cannot control, e.g., sex and age of the donor, storage conditions of the evidence before collection, age of the fingermark, or nature of the substrate, it is necessary to target compounds that are most likely to be found in sufficient quantities in almost all latent fingermarks. Numerous techniques have been developed that target organic and inorganic components with more or less success. Each technique generally is limited in its application according to the type of the substrate and the nature of the secretion. For example, 1,2-indanedione and ninhydrin are applied in the case of latent fingermarks on porous surfaces, cyanoacrylate fuming followed by the application of stains is the most efficient approach for
nonporous ones, physical developer and oil red O are two techniques for detecting
fingermarks when a substrate has been wet, acid yellow 7 and acid violet 17 are specific for
blood-contaminated fingermarks, just to cite a few of several situations encountered. These
limitations find their origin in the intrinsic interaction mechanisms between the reagents and
the secretion residue.

We do not intend to review extensively all the existing fingermark detection
techniques. We shall instead provide a broad overview of the historical and routinely used
methods as well as the techniques for development. Issues concerning application protocols,
formulations, or comparative studies will not be covered. In the context of this review, we
shall classify the techniques according to the kind of stain that is used. The reader must
appreciate that we have adopted a very broad definition of the term, "stain." We define a stain
as any organic molecule, chemical reagent or nanocomposite that permits the visualization of
initially latent fingermarks through the creation of a visible contrast between the ridges and
the underlying substrate. Note that this view is in keeping with definitions existing in the
biological staining literature, such as that of Horobin (1988). The coloration of the secretion
residue can be obtained through 1) physical incorporation of stains in the residue itself, e.g.,
using a lysochrome, 2) chemical reaction between an initially colorless reagent and some
components of the secretion residue, e.g., the amino acids, or 3) physico-chemical processes
leading to specific targeting of the residue, e.g., negatively charged metal nanoparticles, and
its subsequent visualization. Consequently, we begin with stains that color the fingerprints
without undergoing a chemical reaction, continue with stains that chemically react with some
components of the secretion residue, and finally, describe techniques based on nanostructured
materials, a growing field in forensic research.
Physical detection mechanisms

By physical detection, we mean colored molecules or particles that have a physical affinity for the fingermarks and permit their visualization without modification of their molecular structure (Voss-de Haan 2006).

Before addressing the issue of stains, it is worth mentioning that it often is possible to detect fingermarks before any treatment of the evidence by using only ambient light and varying the incident angle. Different optical techniques such as episcopic coaxial illumination (Reiss 1903, Pfister 1985, Ziv and Springer 1993), filtered light (Champod et al. 2004), or reflected ultraviolet (UV) light, i.e., the RUVIS system (Saferstein and Graf 2001) can be used to highlight marks on the substrates. Alternatively, some components of the secretions may show an intrinsic luminescence that can be visualized under UV light (Saitoh and Akiba 2005, 2006) or laser illumination (Dalrymple et al. 1977, Menzel 1999). Most of the time, these detection methods suffice when fresh fingermarks, rich in secretion, are laid on nonporous substrates. If the marks are weak, poor in secretion, old, or on a colored substrate, optical techniques must be extended by using stains.

Powder dusting

Used since the beginning of the last century, powder dusting (Reiss 1911) is one of the first techniques used to detect fingermarks, and it still is widely applied at crime scenes. The principle is simple: powder is dusted onto items using a brush so that it physically adheres to the moist, sticky and greasy substances of fingermark residue (Lee and Gaensslen 2001, Champod et al. 2004). As a result, the contrast between the substrate and the dusted ridges permits observation of the latent fingermarks. This technique is inexpensive and easy to set up, but is limited to smooth and nonporous surfaces. Moreover, only fresh fingermarks give
satisfactory results; the quality decreases as the fingermarks age owing to a loss of stickiness of their residues, which can be problematic.

Conventional powders generally are composed of a contrasting agent, e.g., metal oxides or chromophoric dyes, plus resinous materials, e.g., stearic acid, for adhesion (Sodhi and Kaur 2001). The first colored dusting powders were obtained by using different metal oxides, sulfides and carbonates, e.g., lead, mercury or antimony. These compounds understandably were abandoned because of toxicity issues. Among the routinely used powders currently, a black one is generally applied on light colored substrates. The black color is obtained by including graphite, powdered charcoal, lamp black or molybdenum powder in their composition. On dark backgrounds, white powders, e.g., titanium dioxide, zinc oxide or zinc sulfide, are preferred (Goode and Morris 1983). Another widely used powder, regarded as one of the most effective (James et al. 1991a), is the so called "argentoratum" (Thomas 1973, 1975) composed of flat aluminium particles and 3-5% (w/w) stearic acid. Another commonly used preparation is a gold colored powder composed of bronze flakes. As their name indicates, magnetic powders contain a magnetic carrier in addition to a stain. They require the use of a magnetic brush for dusting, thus avoiding destructive contacts between brush hairs and the secretion residue (James et al. 1991b, 1992, 1993). Luminescent powders are used mainly on multicolored backgrounds, when poor contrast might be expected with conventional powders. A great variety of luminescent dyes or pigments are available, e.g., acridine yellow, coumarin 6, or rhodamine 6G (Lee and Gaensslen 2001). Cyano blue dye, Nile red and proflavine also were tested recently as fluorescent dusting components (Sodhi and Kaur 2004, 2005, Kaur 2006).

An alternative way of applying powders is as suspensions in aqueous solution, a method also known as "wet powdering" or "small particle reagent." This technique will be described below.
**Camphor**

Camphor dusting may be considered an unconventional dusting method, because it requires the use of soot freshly obtained by burning it (Waldoch 1993). Camphor is a flammable white crystalline solid that produces a large amount of black soot when burned. The item to be "dusted" is maintained above the burning camphor to allow the black smoke to cover the entire surface. The soot adheres selectively to the fingermark residues and the excess is removed with a brush or under tap water; the fingermark appears black. Camphor may be helpful for some substrates such as firearms cartridges (Sturelle et al. 2006), but is not used for conventional surfaces.

**Iodine fuming**

Iodine (I$_2$) fuming was described very early (Coulier 1863) for detecting forgeries and fingermarks. A substrate bearing latent fingermarks is exposed to the purple iodine vapors provided by crystals of iodine contained in a glass vessel through which an air stream is forced to pass. The lipids present in the secretions physically absorb the vapors, which results in a temporary yellowish or brownish staining of the marks. Visualization is temporary due to the re-sublimation of iodine vapors from the marks to the ambient atmosphere after a few minutes. It is possible, however, to fix the stained fingermarks by a chemical reaction between iodine and 7,8-benzoflavone (α-naphthoflavone) (Mashito and Makoto 1977, Haque et al. 1983), which results in a permanent blue image. The mark also can be lifted by using the corrosive nature of the iodine. A highly polished silver or tin plate is pressed firmly on the mark for 5 sec, then exposed to a strong white light source or UV lamp. This leads to the formation of a dark image on the plate (Adcock 1977, Arndt 1985).

This technique can be applied to various porous and nonporous surfaces, especially on difficult ones like leather or human skin (Adcock 1977, Gray 1978). It should be emphasized
that only fresh fingermarks can be observed by using this technique. It should also be
mentioned that iodine vapors are toxic and corrosive.

**Lipid dyes: oil red O and solvent black 3**

The lipid fraction of secretion residues constitutes a good target for fingermark detection,
because it is possible to dye them specifically using lysochrome molecules currently used in
biology. Moreover, in the case of fingermarks that have been wet, it is generally accepted that
the water soluble components, e.g., ions, amino acids or proteins, are no longer present on the
substrate due to their solubilization, while the water insoluble components, e.g., lipids, remain
and can still be retrieved.

Oil red O (ORO; solvent red 27) is a relatively new technique for detecting
fingermarks on porous surfaces (Beaudoin 2004, Rawji and Beaudoin 2006, Guigui and
Beaudoin 2007). ORO is a lysochrome used in biology to stain lipids and lipoproteins. When
a porous substrate bearing latent fingermarks is immersed in the ORO working solution, the
fingermarks appear as visible red marks by accumulation of the dye molecules, whereas the
substrate generally undergoes a slight coloration (light red) leading to sufficient contrast
between the mark and the substrate to permit good visualization (Fig. 1). ORO also is used in
forensic science to stain lip marks, especially when lipstick is used (Navarro et al. 2006).

Solvent black 3 (or Sudan black) is another lysochrome used to stain lipids and
phospholipids in histochemistry. Very few applications of this dye for fingerprint detection
have been reported (Hart 2003, Home Office Scientific Development Branch 2005). Solvent
black 3 is applied mostly on greasy, contaminated surfaces for which other methods give poor
results or fail to detect fingermarks. The marks that are obtained after staining are dark
blue.
Enhancement of cyanoacrylate-developed fingermarks

Cyanoacrylate fuming cannot really be considered a physical staining method as defined in the introduction, because chemical processes take place with the secretion residue. This detection technique, however, generally is combined with a subsequent dyeing step, which does fall into this category.

Cyanoacrylate fuming, a method used worldwide, was developed toward the end of the 1970's (Lee and Gaensslen 1984) to detect fingermarks on nonporous surfaces. It involves the use of a colorless monomeric liquid, i.e., the cyanoacrylate ester (also known as Superglue, a rapid bonding, high strength glue) heated to its boiling point. Monomeric vapors are produced that react with some components of fingermarks by polymerization (Fig. 2), but not with the underlying substrate. The exact initiators of the polymerization are not yet clearly determined, because cyanoacrylate shows particular affinity for moisture, lipids and some water soluble components (Lewis et al. 2001, Wargacki et al. 2007). As a result of the polymerization process, which continues until the source of monomers is removed or the heating process is stopped, a white solid covers the developed fingermarks (Mankidy et al. 2006). At the microscopic level, polycyanoacrylate is not a massive and continuous film that covers the secretion residue, but rather can be described as tangled noodles. Owing to the toxicity of the vapors (Hughes 1993), treatment is performed in a cabinet where humidity and heating of the cyanoacrylate can be controlled precisely.

Cyanoacrylate fuming is a sensitive technique for detecting fingermarks on nonporous surfaces. The contrast obtained, however, sometimes is poor due to the white color of the polymer (Fig. 3), which results in barely visible or undetected fingermarks. Fortunately, cyanoacrylate-developed fingermarks can be stained using organic dyes that are trapped selectively in the polymer network and this significantly enhances the visual contrast (Fig. 3).
(Champod et al. 2004). Dyes, typically luminescent ones, that are used routinely in forensic laboratories around the world (Fig. 4) include Ardrox, basic red 28, basic yellow 40, Nile red and rhodamine 6G (McCarthy 1990, Olenik 1992, Day and Bowker 1996). The choice of dye relies mostly on the properties of the substrate, e.g., color, background luminescence, or solvent interactions that govern the specific excitation and emission wavelength domains (Mazzella and Lennard 1995). Solutions of mixed dyes (Olenik 1997), or even rare earth compounds, such as Europium complex, also can be used to widen the wavelength domains (Lock et al. 1995).

Chemical detection techniques

By chemical detection, we mean generally uncolored staining reagents that undergo chemical reaction, i.e., modifications of their molecular structure, with some components of the fingermarks to produce colored or luminescent compounds.

Silver nitrate

A reagent that must be cited for historical reasons is silver nitrate (AgNO₃). This was first used in 1878 to detect chloride ions contained in papillary secretions (Aubert 1877-1878, Olsen 1978). This method can be applied on porous surfaces including paper and untreated wood. The principle is simple: silver nitrate reacts with the chloride ions contained in the secretion residue to form silver chloride (AgCl). Upon exposure to light, AgCl decomposes to form metallic silver (Ag), resulting in a black fingermark. Owing to the diffusion of chloride ions through the substrate, however, blurred fingermarks are obtained if they are aged more than one week. The kinetics of the diffusion has been studied to try to determine the age of the fingermark (Angst 1962).
Amino acid reagents

Amino acids constitute a major organic component of sweat; 0.3–2.59 mg of amino acids are secreted per liter of sweat (Ramotowski 2001), which corresponds to 2.4–20.7 µM for a typical amino acid. Amino acids, therefore, represent a specific target for a fingerprint detection strategy based on chemical reaction. Amino acid detection (or quantification) agents used in biology, such as ninhydrin, have been applied successfully in forensic science as described below.

Amino acids generally, unlike chloride ions, show strong affinity for paper fibers, and their migration through such substrates is prevented. Fingermarks thus are well preserved in the paper matrix and can be detected satisfactorily tens of years after deposition. Note also that amino acids typically are water soluble constituents of sweat. This means that if the item has been wet, such compounds will not be recovered and the following techniques could not be successfully applied.

Ninhydrin (2,2-dihydroxy-1,3-indanedione) was the first amino acid reagent used for detecting latent fingermarks on porous surfaces such as paper and cardboard (Odén and Von Hofsten 1954) and it still is widely used. Ninhydrin can detect primary and secondary amines contained in amino acids or in polypeptides present in latent fingermarks. When reacting in situ with such free amines, ninhydrin produces a deep blue or purple product, Ruhemann's purple (Fig. 5). As a result, fingermarks appear as purple ridges on unmodified substrate (Fig. 6). Several studies have been performed to improve the sensitivity and to optimize the development conditions over wider temperature and relative humidity ranges (Lamothe and McCormick 1972, Grigg et al. 1986, Lennard et al. 1986, Almog 2001, Petraco et al. 2006).

Despite the great sensitivity of ninhydrin, the contrast between the fingerprint and the substrate can be weak, e.g., on dark backgrounds. In this case, observation of the fingerprints is impossible, because Ruhemmann's purple is not luminescent. Efforts have been made to
overcome this problem, especially by using post-ninhydrin treatments containing metal salts. When Ruhemann's purple is reacted with zinc or cadmium ions, luminescent coordination complexes are formed. As a result, fingerprints undergo a color change and become orange instead of purple with zinc and red instead of purple with cadmium. The derivatives also exhibit luminescence when cooled to 77 K with liquid nitrogen (Kobus et al. 1983, Stoilovic et al. 1986, Lennard et al. 1987). Cadmium toxicity and the inconvenience of storing liquid nitrogen, however, are drawbacks limiting application of such post-treatments. Extensive research has been carried out to discover ninhydrin-like reagents with increased sensitivity and room temperature luminescence (Almog et al. 1982, 1992, 2000, 2001, Lennard et al. 1986, 1988, Joullié et al. 1991, Hansen and Joullié 2005). Many ninhydrin analogues have been synthesized and studied, but despite this, few have been used routinely for reasons of expense and time-consuming synthesis protocols.

Another amino acid reagent that has been used widely for nearly 20 years is 1,8-diazafluoren-9-one (DFO) (Wilkinson 2000b). Unlike ninhydrin, DFO produces very faint pink fingermarks, barely visible to the naked eye, but which are extremely luminescent at room temperature (Grigg et al. 1990, McComiskey 1990, Pounds et al. 1990).

More recently, a third amino acid reagent, 1,2-indanedione, has been shown to be effective for detecting fingermarks on porous surfaces. When using 1,2-indanedione, pink colored fingermarks characterized by strong luminescence at room temperature are obtained (Fig. 7). Extensive comparative studies showed that 1,2-indanedione is an alternative to DFO (Joullié et al. 1998, Roux et al. 2000, Wilkinson 2000a, Wiesner et al. 2001, Kasper et al. 2002, Gardner and Hewlett 2003). Recent research also has shown that when zinc chloride (ZnCl$_2$) is introduced to the 1,2-indanedione working solution before treating the specimen, the luminescence of the detected fingermarks is markedly improved (Stoilovic et al. 2007, Wallace-Kunkel et al. 2007, Bicknell and Ramotowski 2008, Russell et al. 2008). For these
reasons and others, such as faster detection procedure, DFO is being replaced by 1,2-
indanedione in the forensic laboratory for detecting fingerprints on porous surfaces.

Concerning the mechanisms of the reactions between amino acids and the reagents
above, a recent study showed that the reactions between DFO or ninhydrin and all the amino
acids go to completion (Spindler et al. 2009). This is not the case with 1,2-indanedione,
however, which varies according to the amino acid involved. The addition of a small amount
of zinc to the working solution seems to play a role in the completeness of the reaction,
because zinc acts as a catalyst.

Two other less widely used reagents for amino acids also are used. Genipin is a natural
product extracted from *Gardenia jasminoides*. Upon reacting with amino acids, a blue
reaction product is obtained that gives a red luminescence at room temperature (Lee et al.
2003, Almog et al. 2004, Levinton-Shamuilov et al. 2005). This reagent constitutes the only
"dual fingermark reagent" for amino acids, i.e., a reagent giving visible/colored fingermarks
that also can be observed with luminescence. Genipin has the further advantage of being a
natural and nontoxic product. Lawsone (2-hydroxy-1,4-naphthoquinone) is the compound
thought to be responsible for the staining properties of henna, commonly used as a skin and
hair dye. In forensic science, naphthoquinones represent a new group of compounds that can
be used to detect the amino acids (Jelly et al. 2008). After treatment with lawsone,
fingermarks are stained purple-brown and exhibit a strong red luminescence. Currently, only
experimental results are available and further research is needed before considering it an
effective reagent.

*p-4-Dimethylaminocinnamaldehyde (pDMAC)*

Urea is another component of sweat that could be targeted to detect fingerprints on porous
surfaces. The first reagent proposed for this purpose was p-4-dimethylaminocinnamaldehyde
(pDMAC). The resulting fingermarks are red and luminescent (Sasson and Almog 1978, Brennan et al. 1995, Ramotowski 1996). The sensitivity of pDMAC, however, has been shown to be less than the conventional amino acid reagents described above. Moreover, blurred fingermarks generally are obtained if they are more than two days old. Finally, the actual molecular target of pDMAC has been questioned (Home Office Scientific Development Branch 2006) and it appears that pDMAC actually could be reacting with amino acids rather than with urea. pDMAC is not used routinely.

**Osmium tetroxide and ruthenium tetroxide**

Osmium tetroxide (OsO$_4$) and ruthenium tetroxide (RuO$_4$, RTX) are compounds that can react with the double bonds of the unsaturated organic compounds found in a fingermark deposit through an oxidation mechanism (Grzegorzewska and Filbrandt 2004). Direct exposure to OsO$_4$ vapors provides black fingermarks after 1 to 12 h (Olsen 1978), whereas RTX gives gray marks after only 10 to 20 min (Mashiko et al. 1991, Mashiko and Miyamoto 1998). Despite the excellent results that can be obtained on a variety of surfaces, especially the problematic ones like human skin, RTX and OsO$_4$ are not used routinely because of their high toxicity through inhalation or skin contact and hazards associated with their manipulation (Blackledge 1998). Only fully equipped laboratories can consider their use.

**Fingermarks in blood**

When the victim and/or the offender are injured, hands can become contaminated with blood. In such cases, bloody fingermarks are likely to be recovered. Depending on the amount of blood and on the nature of the substrate, some marks are obvious to the naked eye, but others remain latent and require the application of specific blood reagents.
Although two different categories of blood reagents exist, viz., protein stains and heme reagents, blood sometimes can be visualized directly due to a strong absorption at 415 nm (Stoilovic 1991) that makes it appear black under illumination at this wavelength. Blood also can be observed due to luminescence in the long-wave UV region when short-wave UV exciting light is used (Springer et al. 1994). Before using any blood-specific reagent, the items bearing latent fingermarks generally are dipped in methanol or in an aqueous 5-sulfosalicylic acid solution (2% w/v) to avoid loss or diffusion of blood due to the water-based blood reagents (Hussain and Pounds 1988, Sears and Prizeman 2000). Once fixed, effective blood reagents can be applied.

Protein stains are compounds adopted directly from biology. The application protocol consists of immersing the item to be treated in a dye solution. The staining times vary from 10 sec to several minutes depending on the dye, the type of substrate and the amount of blood. The dye binds to the proteins, which leads to a colored or luminescent complex. The item then is washed with successive de-staining solutions, based on methanol or water, to remove the excess dye and to clear the background so that contrast is improved (Fig. 8). Because numerous protein stains can be applied, only the commonly used ones are noted here (Sears and Prizeman 2000, Sears et al. 2001, 2005, Marchant and Tague 2007). A conventional non-luminescent stain is amido black (naphthol blue black B), which gives dark blue marks on light blue or nearly colorless substrates. Luminescent stains also are available, e.g., acid yellow 7, acid fuchsine (Hungarian red) and acid violet 17 (Coomassie brilliant violet). Which dye is used depends on the color and the nature of the substrate.

Heme reagents are molecules that can undergo oxidation, initiated by hydrogen peroxide in presence of a catalyst, in this case the heme group of hemoglobin. As a result, colored, luminescent, or chemiluminescent (a product generating its own light with no need to use excitation light source) products are formed. The reaction is selective for blood (possible
false positive results are not discussed here) and gives little background staining. Routinely
used stains include 3,3-diaminobenzidine (DAB) (Allman and Pounds 1991, 1992, Sahs
1992), which gives a dark brown color; ABTS (2,2'-azino-di-(3-ethyl-
benzthiazolinesulfonate) diammonium salt) (Caldwell et al. 2000), which gives a green
reaction product; and leucomalachite green, phenolphthalein, leucocrystal violet, o-tolidine
(Lee 1984), fluorescein (Cheeseman and DiMeo 1995), leuco rhodamine 6G (Yapping and
Yue 2004) or eosin Y (Wang et al. 2007). Another heme reagent that must be cited, even if it
is not used as a fingermark reagent, is luminol. This gives rise to intense blue
chemiluminescence and is used to detect traces of blood as might be found on a crime scene
or on items that have been washed (Webb et al. 2006). Any fingermarks that are obtained,
however, are not sufficiently sharp to detect fingerprint ridges that are necessary for
identification of the person, and the procedure is used only to detect the presence of blood,
which facilitates subsequent blood sampling for DNA analysis.

Finally, note that the conventional fingermark reagents, e.g., amino acid reagents or
cyanoacrylate, also can detect blood fingermarks even if they do not possess the sensitivity of
the specific blood reagents. Conventional stains, therefore, generally are applied before the
blood reagents to detect most of the existing bloody or non-bloody fingermarks.

Nanoparticles – a new generation of stains

During the last decade, considerable interest has emerged within the forensic science
community in using nanoparticles or nanocomposites for detecting fingermarks.
Nanoparticles already are applied successfully in cell and whole animal imaging (Bagwe et al.
2007), but constitute an emerging field in forensic science.
Interest in nanoparticles to detect fingermarks is due to their properties: 1) their size is 1,000 to 10,000 times smaller than a fingerprint ridge width, which ensures excellent resolution upon detection, 2) their structure can be modified easily by addition of molecular chains or chemical functionalities on their outer surface, and 3) their optical properties make them excellent candidates for obtaining luminescent fingermarks. Nanoparticles such as quantum dots are intrinsically luminescent (Bawendi et al. 1990, Wang and Herron 1991, Murray et al. 1993), and others, e.g., silica nanoparticles, are made luminescent by incorporating organic dyes in their structure during their synthesis.

The challenge for using nanoparticles to detect fingermarks lies in the fact that nanoparticles per se generally have no specific affinity for the marks. It is necessary, therefore, to find a way to make nanoparticles specifically target the secretion residues without acquiring an affinity for the substrate; otherwise the resulting contrast would be poor (Fig. 9). A successful application of nanoparticles generally is obtained by adding functional groups onto the nanoparticles (Fig. 10) or by modifying the experimental conditions, such as the pH, to influence parameters such as the charge on the nanoparticle.

The following sections describe four approaches that were selected for their specificity in terms of fingermark detection processes: physico-chemical detection, lipid-lipid interaction, and electrostatic interaction.

**Metal nanoparticles**

Use of metal nanoparticles to detect fingermarks on various kinds of substrates has interested forensic scientists for several years (Choi et al. 2008b). Both in terms of research and practical applications, most interest has focused on two kinds of metal nanoparticles, gold and silver. The multimetal deposition technique (based on gold nanoparticles) and physical development (based on silver nanoparticles) are not new; both were developed in the 1980s. Research
continues to clarify understanding of the underlying physico-chemical processes that permit
the detection of fingermarks, however, with the aim of enhancing and simplifying existing
protocols. Both approaches are described below, followed by a section dealing with additional
possibilities offered by metal nanoparticles.

Multimetal deposition (MMD)

Gold nanoparticles conventionally are synthesized in water. A gold (III) derivative (generally,
tetrachloroauric acid; HAuCl$_4$) is reduced using an agent such as sodium citrate or sodium
borohydride. The reduction of HAuCl$_4$ by sodium citrate in boiling water is one of the most
commonly used procedures for obtaining monodisperse spherical gold nanoparticles,
generally about 10–20 nm diameter, with time stabilities of several months to years
(Turkevich et al. 1951, Frens 1973, Slot and Geuze 1981, Turkevich 1985a,b). The origin of
such excellent stability is to be found in the capping of the gold nanoparticles in solution by
tri-negatively charged citrate ions. The electrostatic repulsion of the resulting negatively
charged nanoparticles prevents them from aggregating. The same negative charge also is the
basis of their ability to detect fingermarks by exploiting charges on organic components such
as proteins, amino acids and lipids as latent prints deposited on a substrate.

If a sample bearing a fingermark is immersed in an aqueous solution of freshly
prepared colloidal gold (pH ~ 6.2), nothing happens. If the pH of the colloidal solution is
lowered beforehand, however, it is possible to find a small range of pH for which both of the
following conditions are met. 1) The tri-negatively charged citrate ions are not yet fully
protonated (pKa$_1$ = 3.13, pKa$_2$ = 4.76, pKa$_3$ = 6.40) and still give the gold nanoparticles a
negative charge. It should be noted that the interaction of carboxylic groups with a metal
surface, as is the case with sodium citrate and gold, can lead to a decrease of the pKa values
(increasing acid strength of the groups) measured for the same molecule in solution.
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(Królikowska and Bukowska 2007). 2) The proteins and amino acids that are entrapped in the residue become progressively more positively charged due to the protonated amino groups (-NH$_3^+$). These positive charges are progressively less counterbalanced by the carboxylate groups (-COO$^-$) as they become protonated to neutral carboxylic moieties (-COOH) at low pH.

In the case of fingermarks, the range of pH is extremely narrow and has been determined experimentally as 2.5-2.8 (Schnetz and Margot 2001). When the pH of colloidal gold is within this range and a sample bearing a fingermark is dipped in it, negatively charged gold nanoparticles in solution become attracted electrostatically by the positively charged residue. Above this range, proteins are not sufficiently positively charged; below this range, gold nanoparticles become uncharged by complete protonation of the sodium citrate ions (the solution turns dark purple as the gold nanoparticles aggregate owing to lack of electrostatic repulsion). This mechanism of staining selectivity is intriguingly like that of acid dyeing of proteins in biological staining (Horobin 1988) and textile dyeing (Zollinger 2003).

These physico-chemical considerations are the basis of the MMD technique, (Fig. 11). MMD was developed in 1989 (Saunders 1989) and improved in 2001 (Schnetz and Margot 2001). This technique is based on the electrostatic attraction between the gold nanoparticles and secretions when the pH of the colloidal gold is adjusted to 2.65. Fingermarks detected by gold nanoparticles only, however, are not readily visible to the naked eye, because they produce very faint pink marks due to the small size of the nanoparticles. To increase the contrast, a second metal deposition is performed by reducing silver ions to metallic silver on the surface of the gold nanoparticles, which play the role of catalytic sites. As a result, fingermarks appear dark brown as illustrated in Fig. 12. MMD is an efficient technique for a wide range of substrates, e.g., porous, nonporous and wet, but it is highly labor intensive and produces dark brown marks only.
An alternative to MMD was proposed in 2007, the single metal deposition (SMD) (Stauffer et al. 2007, Durussel et al. 2009). The procedure is similar to MMD, except that the enhancement step, i.e., second metal deposition, is performed by reducing gold ions instead of silver onto the gold nanoparticles. The results obtained with SMD are similar to MMD (Fig. 12), but the overall procedure is cheaper, less labor intensive, and easier to set up.

Both MMD and SMD produce dark brown fingermarks. This means that the contrast can be very good on transparent, white or light colored substrates, but these techniques fail to produce sufficient contrast on dark colored or patterned substrates for which purpose luminescence is required. A modification of the enhancement step led to the formation of a zinc oxide shell instead of silver or gold around the gold nanoparticles after they were deposited on the secretion residue (Becue et al. 2008). In this way, luminescent fingermarks were obtained on dark, nonporous substrates (Fig. 12). This technique currently is limited to nonporous surfaces, however, unlike MMD and SMD, which can be applied on porous and nonporous substrates.

Physical developer

Use of a physical developer (PD) is another detection technique based on metal nanoparticles and driven by physico-chemical mechanisms (Goode and Morris 1983). Unlike MMD, for which metal colloids were synthesized prior to the development procedure, PD consists of immersing samples bearing fingermarks in an acidic aqueous solution containing silver ions e.g., silver nitrate; reducing agents, e.g., ferrous and ferric ions; positively charged ionic surfactant, e.g., n-dodecylamine acetate; and other chemical components such as citric acid to lower the pH of the solution. The accepted mechanism is that during the reduction process, negatively charged silver colloids are formed by reduction of the silver ions (Cantu 2001, Cantu and Johnson 2001). If such silver nanoparticles are formed in solution, they are
surrounded rapidly by positively charged surfactant molecules, which prevent silver colloids from aggregating and stops particle growth (Fig. 13). Such nonreactive nanoparticles do not play a role in fingermark detection and remain in solution. If the negatively charged silver colloids are formed near a fingermark, however, they immediately become attracted by the positively charged residue because the pH of the solution is lowered to the acidic range. Once entrapped in the secretion, these silver colloids act as reduction sites for the additional silver ions. As a result, fingermarks become progressively visible as the reduction process continues. At some point, the forensic scientist stops the process by removing the sample from the reducing bath and by dipping it in a neutralizing/rinsing water bath. The fingermarks obtained are a gray color (Fig. 13). Like MMD, PD is applied to white colored substrates only.

Other applications of metal nanoparticles

In the cases of MMD, SMD and PD, metal nanoparticles are used without further surface modifications; the detection is driven by physico-chemical mechanisms and especially electrostatic interactions. Another strategy involves functionalizing the metal nanoparticles with nonpolar hydrophobic aliphatic chains (Fig. 14) so that they can be applied in suspension in organic solvents (Sametband et al. 2007) or as enhanced nanopowders once dried and dusted using a brush (Choi et al. 2006, 2008a). The addition of aliphatic chains to the nanoparticles enhances the interactions between the metal nanoparticles and the lipids contained in the fingermark residue (Choi et al. 2008b). Compared to classical (micro-sized) powders, nanopowders have been shown to give better results in terms of ridge details and less unwanted deposition on the underlying substrate. Despite this, the use of nanopowders did not necessarily lead to a better contrast, because the detected fingermarks were a lighter color than the dark colored classical powders.
Another research strategy consisted of binding antibodies onto gold nanoparticles to take advantage of the highly selective antibody-antigen recognition process to detect specific drug metabolites in a fingermark (Leggett et al. 2007). In an attempt to provide evidence of the consumption of a drug in addition to touching contaminated objects, anti-cotinine antibodies bound to gold nanoparticles have been used to target cotinine, a metabolite of nicotine, present in the sweat of tobacco smokers. Combined with a fluorescent marker, highly detailed fingermarks can be observed on glass slides (Fig. 15). Another research strategy is immunolabeling to identify antigenic sites with the secretion residue components or keratinized skin. Proteins present in sweat or skin remnants that have been identified in this way are keratins 1 and 10, cathepsin D and dermcidin using the corresponding antibodies (Drapel et al. 2009).

**Metal Oxide nanoparticles**

*Nanopowder and small particle reagents*

Metal oxides have been widely used for detecting fingermarks due to their optical properties, i.e., color and fluorescence, because it is possible to obtain both white particles, such as titanium dioxide (TiO$_2$) or zinc oxide (ZnO), and also black particles, such as iron oxide (Fe$_2$O$_3$). Such reagents can be used as conventional fingerprint powders or they can be suspended in aqueous media using a detergent to enhance their solubility. As a result, one can detect fingermarks on nonporous substrates or on the adhesive side of tapes using a process that has been described as a "wet version of the classical powdering process" or "wet powdering" (Cucè et al. 2004, Williams and Elliot 2005). Instead of using a brush, samples are treated by immersing them in particle suspensions before rinsing with water. Fingermarks are detected by entrapment of the metal oxide particles in the lipid matrix of the secretion residue. This technique uses "small particle reagents" (SPR) and gives good results with
nonporous substrates on which latent fingermarks are present, especially if the substrate has been wet (Goode and Morris 1983, Frank and Almog 1993, Champod et al. 2004). In addition to the metal oxide particles noted above, black molybdenum sulfide (MoS$_2$) and white zinc carbonate (ZnCO$_3$) particles also are used as SPR agents. According to the color of the underlying substrate, white powder, e.g., TiO$_2$, or black powder, e.g., Fe$_2$O$_3$, is used to maximize the contrast. A luminescent version of SPR has been proposed that involves addition of luminescent organic dyes, e.g., basic yellow 40 in ethanol, to a conventional aqueous SPR solution (Springer and Bergman 1995, Lee and Gaensslen 2001, Jasuja et al. 2008).

The size of the particles influences the result of SPR; smaller particles give better results than larger ones (Frank and Almog 1993). As for dried nanopowders, the use of nanoparticles in the context of SPR can greatly improve the resolution.

More recently, nanostructured ZnO was used to detect fingermarks on nonporous surfaces by taking advantage of its visible fluorescence when dry dusted or applied as an SPR (Choi et al. 2008a). These investigators combined oleylamine with the fluorescent dye, perylene dianhydride, to form an entity that then was adsorbed onto TiO$_2$ nanoparticles to form a powder exhibiting strong fluorescence at 650-700 nm when excited at 505 nm (Choi et al. 2007). When dusted onto nonporous substrates bearing latent fingermarks, this fluorescent nanopowder was slightly weaker in fluorescence intensity compared to conventional luminescent powders, but produced significantly less background. As a result, good contrast was obtained between the fingermarks and the substrate.

*Titanium dioxide as a blood reagent*
Sub-micron titanium dioxide particles suspended in anhydrous methanol can be used to detect blood fingermarks on nonporous and "semiporous" surfaces (Bergeron 2003). The results were excellent on nonporous surfaces. No mechanistic explanation was offered for the affinity of TiO$_2$ for blood. Uptake, however, may be related to the fact that TiO$_2$ has some affinity for blood (Nygren et al. 1997) and for proteins by electrostatic interactions (Topoglidis et al. 2001).

**Quantum dots (QDs)**

QDs are crystalline semiconductor nanoparticles containing a few hundred to several thousand atoms varying in size from 1 to 10 nm. Owing to quantum confinement, QDs exhibit strong luminescence at room temperature. It is of interest that the emission wavelength is dependent on the size of the nanoparticle; larger nanoparticles emit in the infrared/red range and smaller ones emit in the ultraviolet/blue range of the spectrum. Another significant feature of such nanoparticles is the fact that all the QDs have a broad excitation spectrum in the ultraviolet range. This means that a variety of QDs, each emitting its own wavelength, can be excited by a single excitation source. These nanoparticles are used in a wide variety of applications such as components for lasers (Bukowski and Simmons 2002) or as fluorescent probes for cellular imaging (Medintz et al. 2005). QDs high quantum yield and ability to resist photobleaching make them superior markers compared to organic fluorescent dyes (Resch-Genger et al. 2008). QDs can be synthesized in either water or organic solvents depending on their external surface coating. Moreover, their outer surface can be functionalized easily with chemical groups to provide new functionalities. Consequently, QDs are widely used in cell biology to stain different components of cells or tissues (Bruchez et al. 1998).

In forensic sciences, particularly for fingermark detection, the properties of QDs are stimulating great interest. Their surfaces can be coated specifically to target the secretion
residue and the luminescence emission color can be tuned in case of a luminescent background. Since 2000, several studies have focused on the use of QDs to detect fingermarks. Cadmium sulfide (CdS) nanocrystals, used alone or combined with dendrimers, i.e., hyperbranched polymers, have been proposed as stains for cyanoacrylate-developed fingermarks (Bouldin et al. 2000, Menzel et al. 2000a,b). The results, however, were disappointing, because the staining times required were too long (24 h) and the working solutions were too unstable. More recent studies have provided only limited improvements (Yu-Juan et al. 2008). The combination of QDs and dendrimers does not seem to constitute an alternative to the existing cyanoacrylate stains.

In another recent study, researchers stabilized QDs in petroleum ether by grafting aliphatic chains on their outer surface and attempted to detect sebaceous fingermarks with these nanomaterials. Promising results were obtained on silicon wafers, but on porous surfaces, the results were limited due to high background luminescence (Sametband et al. 2007).

Cadmium sulfide (CdS) encapsulated in a biopolymeric chitosan matrix has been used as dusting powder for detecting fresh marks on aluminium foil (Dilag et al. 2009). Luminescent fingermarks were obtained, but the process involved in the detection is only physical. Moreover, the use of dried cadmium-based nanoparticles as dusting powder has serious health and safety implications. Water soluble cadmium selenide (CdSe) QDs have been synthesized and used to detect fresh marks on the adhesive surface of tape with promising results (Wang et al. 2009). Cadmium telluride (CdTe) QDs, synthesized in aqueous solution, were used to detect blood fingermarks on various nonporous surfaces (Becue et al. 2009) such as aluminium foil, black polyethylene, glass or transparent polypropylene (Fig. 16). A comparison with acid yellow 7, one of the best blood reagents for nonporous substrates
(Sears et al. 2001), showed that the QDs were superior to acid yellow 7 on aluminium and equally effective for the other substrates.

Currently, the use of QDs to detect fingermarks remains a research proposition and no casework uses of QDs have been reported. QDs show great promise, however, for efficient fingermark detection. Issues remaining to be addressed before routine application is likely include the toxicity of cadmium and improved targeting of secretion residue by functionalization of the QD outer surface.

**Silica nanoparticles**

Synthesis of silica nanoparticles is simple and consists of a succession of hydrolysis/condensation processes on silicon alkoxide precursors (Si-(O-R)_4; R = an alkyl chain). This leads to the formation of silicon oxide nanoparticles in suspension in water (Stöber and Fink 1968) or in a water-in-oil emulsion (Arriagada and Osseo-Asare 1999, Bagwe et al. 2004). The introduction of organic dyes during the synthesis permits their attachment to the nanoparticle surface or to their embedment within the particles (van Blaaderen and Vrij 1992, Santra et al. 2001, Tapec et al. 2002, Zhao et al. 2004, Johnston et al. 2005). Such dye-doped nanoparticles currently are widely used as labeling agents for bio-imaging purposes. They are highly efficient alternatives to classical organic fluorophores, because the silica coating (or encapsulation) of some dyes isolates them from the oxygen and water of the outside environment. This both increases photostability and emission quantum yield of the dye and decreases photobleaching by preventing penetration of oxygen, which promotes photodecomposition (Tan et al. 2004).

In the context of fingermark detection, the use of luminescent silica nanoparticles for forensic purposes remains at the pilot study stage. Theaker et al. (2008) recently reported entrapment of a variety of colored and fluorescent dyes including basic red 28, basic yellow
40, fluorescein, methylene blue, oxazine perchlorate, rhodamine B, rhodamine 6G and thiazole orange within silica particles. The resulting doped nanoparticles were used in aqueous suspensions to detect fingerprints. The process is very similar to that described for small particle reagents. Micron-size particles also were used as dusting agents. Both fresh (20 min) and aged fingerprints (40 days old) showed good definition after development.

Current research projects in our laboratory aim to use dye-doped silica nanoparticles that are functionalized chemically to recognize the fingerprint residue. Such dye-doped nanoparticles can be considered inert labeling agents that require appropriate functionalization to promote (bio)molecular interactions or reactions with components of interest. One advantage of the silica nanoparticles is the versatility provided by their surfaces for chemical modification. This is because the surfaces of silica nanoparticles carry silanol groups (Si-OH). These moieties readily undergo chemical modifications, such as the addition of an additional siloxane layer using an organosilane or functionalized alkoxyisilane, e.g., X-(CH₂)m-Si-(OR)₃; where X is the functionality to be added. Other chemical modifications include addition of alkyl chains bearing amine, carboxyl, or thiol functionalities (Fig. 17). The incorporation of chemical groups onto the silica surface allows tailoring of the chemical behavior of the surface, and thus the nanoparticle as a whole, compared to inert "naked" particles. The challenge in forensic science is to find the best functionalities to add to dye-doped silica nanoparticles to enhance targeting of latent fingerprints on a wide range of substrates.

Preliminary results are encouraging and suggest the considerable potential of such a strategy for detecting fingerprints by combining the intense luminescence of the dye-doped silica nanoparticles and the selectivity of the grafted ligands.

In forensic science, and more particularly in the context of fingerprint detection, stains play a key role. To cover the subject as completely as possible, we chose to define a "stain" as every chemical entity (organic molecule, metal nanoparticle, dye-doped nano- and micro-
structured element, or other compound) that can create a contrast between a latent fingermark and its underlying substrate. This wide definition goes beyond the conventional organic dyes or organic dye-doped nanoparticles that currently are used for bio-imaging applications, because it also encompasses entities such as iodine, metal salts, or metal nano- or micro-particles, which are excellent contrast elements.

With this review, we aimed at presenting various aspects of fingermark detection using dyes. The possibilities offered to forensic scientists are numerous and include conventional staining of the lipid fraction of fingermarks, chemical targeting of amino acids or proteins, selective polymerization on the secretion residue, or the use of more complex physico-chemical processes. The choice for one technique over another, or for a combination of different ones, is determined mainly by the nature of the item or by the surface at the crime scene, e.g., walls or furniture, and by the context of the criminal affair, e.g., timeline, bloody case, evidence exposed to environmental elements, prolonged exposure to water. In addition, the choice for a luminescent dye or a conventional stain is strongly dependant on the nature and color of the substrate, even if it is generally recognized that luminescent techniques are more sensitive compared to nonluminescent ones.

During the last decade, several investigations have focused on the development of new detection techniques based on the use of functionalized and/or luminescent nanoparticles for detecting fingermarks. Nanoparticles have great promise for fingermark detection due to their high surface-to-volume ratio, their size-dependent qualities, their optical properties and the fact that they can easily be tuned chemically. If surface modification and composition control are combined, these properties impart high selectivity and sensitivity. The results currently are promising and could provide forensic scientists with new and efficient alternatives to the currently existing techniques.
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**Fig. 1.** Left: molecular structure of oil red O. Right: representative fingermark detected using this technique with paper as substrate.

**Fig. 2.** Polymerization mechanism on the secretion residue during the cyanoacrylate fuming process.

**Fig. 3.** Left: cyanoacrylate-fumed fingermark located on a beer glass. Note that the quality of development is not optimal. Right: same fingermark after enhancement using rhodamine 6G as a cyanoacrylate dye and observed using luminescence.

**Fig. 4.** Chart illustrating the excitation and emission wavelength ranges for the most commonly used cyanoacrylate dyes, the colors of which are indicated on the diagram.

Abbreviations: BY40, basic yellow 40; BR28 basic red 28; S7 Styryl 7.

**Fig. 5.** Reaction mechanism between ninhydrin and amino acid leading to the formation of a dark purple product, Ruhemann's purple.

**Fig. 6.** Left: molecular structure of ninhydrin (2,2-dihydroxy-1,3-indanedione). Right: representative fingermark detected using this technique with paper as substrate.

**Fig. 7.** Left: molecular structure of 1,2-indanedione. Right: representative fingermark detected using this technique with paper as substrate and observed using luminescence.
**Fig. 8.** Left: molecular representation of acid yellow 7. Right: representative blood fingerprint detected using this technique with aluminium foil as substrate.

**Fig. 9.** Schematic illustration of the main characteristics of nanoparticles influencing their use as fingerprint detection reagents. Good contrast is obtained when the ridges of the mark are targeted specifically, but not the underlying substrate.

**Fig. 10.** Schematic illustration of the functionalization strategy involving addition of ligands on the outer surface of nanoparticles to increase affinity for the secretion residue.

**Fig. 11.** Schematic illustration of the two-step metal deposition application protocols. This chart shows MMD, SMD, and MMD\textsubscript{lumin}, which differ in terms of which metal is used for enhancement, silver, gold, and zinc, respectively.

**Fig. 12.** Schematic representation of the differences among the three metal deposition protocols in terms of final nanoparticle composition and detected fingerprints. MMD and SMD produce dark colored fingerprints, whereas luminescent MMD (MMD\textsubscript{lumin}) generates luminescent fingerprints.

**Fig. 13.** Schematic illustration of the two processes involved in the physical developer detection technique in terms of the distance between the secretion residue and the site of colloid formation. Top: in solution. Bottom: close to the residue.

**Fig. 14.** Left: schematic representation of the oleylamine-functionalized gold nanoparticle used as nanopowder dusting on fingerprints (Choi et al. 2006). Right: schematic
representation of the alkanethiol-capped gold nanoparticle (Au-NP-C18) used to detect sebaceous fingermarks on porous and nonporous surfaces in an enhanced physical developer process (Sametband et al. 2007).

**Fig. 15.** Schematic representation of the "protein A/antibody"-functionalized gold nanoparticle used to detect fingermarks by antibody/antigen recognition. The illustrated fingermark was obtained from a male smoker after 40 min sweating and detected using the anti-cotinine-functionalized nanoparticles with Alexa Fluor 546 as luminescent marker. Fingermark reproduced with permission from (Leggett et al. 2007), Copyright Wiley-VCH Verlag GmbH & Co. KGaA.

**Fig. 16.** Left: schematic representation of the CdTe/CdS quantum dot used to detect fingermarks on nonporous substrates (Becue et al. 2009). Right: representative fingermark detected using this technique with aluminium foil as substrate.

**Fig. 17.** Potential outer shell functionalization of a silica nanoparticle with the objective of promoting the interactions with organic compounds such as amino acids and proteins in the secretion residue.
(Left) Molecular representation of Oil Red O; (Right) Representative fingermark detected using this technique (paper as substrate).

109x61mm (400 x 400 DPI)
Polymerization mechanism taking place on the secretion residue during the cyanoacrylate fuming process.

133x33mm (400 x 400 DPI)
(Left) Cyanoacryle-fumed fingermark located on a beer glass (the quality of development is not optimal), (Right) Same fingermark after enhancement using Rhodamine 6G as cyanoacrylate dye (observation in luminescence).

140x76mm (400 x 400 DPI)
Chart illustrating the excitation and emission wavelength ranges for the most commonly used cyanoacrylate dyes.

99x63mm (300 x 300 DPI)
Reaction mechanism between Ninhydrin and amino acid, leading to the obtaining of a dark-purple product: the Ruhemann's purple.

99x66mm (300 x 300 DPI)
(Left) Molecular representation of Ninhydrin (2,2-dihydroxy-1,3-indanedione); (Right) Representative fingermark detected using this technique (paper as substrate).

109x64mm (400 x 400 DPI)
(Left) Molecular representation of 1,2-Indanedione; (Right) Representative fingermark detected using this technique (paper as substrate; observation in luminescence).
108x64mm (400 x 400 DPI)
(Left) Molecular representation of Acid Yellow 7; (Right) Representative blood fingerprint detected using this technique (aluminium foil as substrate).

120x64mm (400 × 400 DPI)
Schematic illustration of the main characteristics of nanoparticles from which benefit should be taken to be used as fingermark detection reagents. A good contrast is obtained if the ridges are specifically targeted, and not the underlying substrate.

150x118mm (400 x 400 DPI)
Illustration of the functionalization strategy consisting in adding ligands on the outer surface of nanoparticles to increase their chemical potential in terms of affinity for the secretion residue.

200x53mm (400 x 400 DPI)
Simplified illustration of the "two-steps" metal deposition application protocol. This chart is valid for multimetal deposition (MMD), single-metal deposition (SMD), and luminescent MMD (MMDlumin), which differ by the choice of the metal in the enhancement step: silver, gold, and zinc oxide, respectively.

127x39mm (400 x 400 DPI)
Schematic representation of the differences between the three metal deposition protocols in terms of final nanoparticle composition and detected fingermark aspect. Multimetal deposition (MMD) and single metal deposition (SMD) permit to obtained dark-coloured fingermarks, whereas luminescent MMD (MMDlumin) permits the observation of luminescent fingermarks.

149x138mm (400 x 400 DPI)
Illustration of the two processes involved in the physical developer detection technique, according to the distance between the secretion residue and the colloid formation site ("in solution" or "close to the residue").

175x79mm (400 x 400 DPI)
(Left) Schematic representation of the oleylamine-functionalized gold nanoparticle used as nanopowder to be dusted on fingermarks (Choi et al., 2006). (Right) Schematic representation of the alkanethiol-capped gold nanoparticle (Au-NP-C18) used to detect sebaceous fingermarks on porous and non-porous surfaces in an enhanced physical developer process (Sametband et al., 2007).

190x101mm (400 x 400 DPI)
Schematic representation of the "protein A / antibody"-functionalized gold nanoparticle used to detect fingermarks by taking benefit of the highly-specific antibody/antigen recognition process. The illustrated fingermark has been obtained from a male smoker after 40 min sweating and detected using the anti-cotinine-functionalized nanoparticles and Alexa Fluor 546 as luminescent marker (source of the fingermark: (Leggett et al., 2007)). 200x114mm (400 x 400 DPI)
(Left) Schematic representation of the CdTe/CdS quantum dot used to detect fingermarks on non-porous substrates (Becue et al., 2009); (Right) Representative fingermark detected using this technique (aluminium foil as substrate).

123x76mm (400 x 400 DPI)
Potential outer shell functionalization of a silica nanoparticle having for aim to promote the interactions with organic compounds such as amino acids and proteins contained in the secretion residue.

132x52mm (400 x 400 DPI)