Nanocomposite Materials with Oriented Functionalized Structure

Petrişor Zamora Iordache, Nicoleta Petrea, Rodica Mihaela Lungu, Răzvan Petre, Ciprian Său and Ioan Safta
Scientific Research Center for CBRN Defense and Ecology
Romania

1. Introduction

Functionalized materials are material structures able to meet certain functions and tasks in well established interaction conditions. The functionality of a nanostructured material is completely conditioned by the well-defined relation holding between the functionality of the host structure (carrier support - CS) and the structure upon which it is carried out (target structure - TS). The functionality and the functions of a material are carried out by means of its physical, chemical or other characteristics, and by means of which it interacts with neighbour material systems, in order to fulfil the assigned functions. The most well-known examples of functionalities are those of: chemical reticulation, detection, controlled stimulation, chemical separation, etc. (fig.1). Most of the natural material structures are found in the form of nano or micrometrically structured composites, and dependency and interaction relations are established between their structural elements, depending on the internal structure and the physical and chemical characteristics of the neighbouring material structures. Composite materials are mixtures of material phases, either amorphous, or structurally well delimited, varied from a morphostructural and morphochemical point of view. The basic structure of composite materials is represented by nanostructured material phases, as material superior form of organization of atomic and molecular structures. As a result of physical and chemical interactions, the structural components of natural composites establish complex physical and chemical equilibrium states, characterised by the fact that the functional structure of this type of composites is masked or cancelled.

The functionality concept is a relatively defined concept, taking as a reference point the investigation and order needs of the surrounding material structures. Basically, the function of a functionalized material defines the way in which the functional structure affects a certain process or a certain material structure, not necessarily neighbouring it. All material structures are carriers of certain types of inherent functionalities, by means of which they specifically interact with the exterior, but which do not have selectivity for certain target structures. Most modern scientific and technological applications need nanostructured or nanocomposite materials, having an oriented functional structure. These constrains impose that natural composite materials (CM) should be submitted to morphostructural and morphochemical modelling processes, in order to use them as oriented functionality materials. From the point of view of the sense of functionality orientation, the materials with oriented functional structure (M_{OF}) can be defined as materials whose structure and
characteristics model and transform in sequences that are phenomenologically ordered in space and time, the structural evolution and the properties of the target structures (TS).

Fig. 1. Some typical examples of nanostructures with orientated functionality (a) Polyfunctional structure with macromolecular separation tasks (ES – external magnetic field stimulus; CL – coated layer; MNP – magnetic nanoparticles; T – target macromolecules; CLM – reticulation molecules; F – functionalities) (b) Functional structure of a interfaces with sensitization tasks (EI – electric interface; SC – signal carrier; ST – signal transducer; SI – structural interface; PL – protective layer; S – external stimulus)

Fig. 2. Typical diagram of an orientated process for controlled separation of organic pollutants. Figure 2 presents the functional diagram of a M_{OF} intended for the reticulation, degradation and controlled separation of the organic pollutants in the surface waters. The material obtained by means of controlled functionalising processes was modelled so that it could carry out the nano- and microfiltering (F_1), reticulation (F_2), degradation (F_3), magnetic separation (F_4) and biodegradation (F_5) functions of the organic pollutants in the surface waters, containing chemical functionalities of the aminated, hydroxilated, carboxylated, phenolated, carbonylated, thiolated and halogenated types, derivatives or chemical structures having mixed functionality. In phase S_1, the material comes in contact with the pollutants (P), reticulating them at the level of the functional structures of the CL and CS_p structural components of the composite. CL carries out the compositing functions of the CS_p
and NP$_S$ structural elements (fig.2) and the filtering functions at molecular level of the contaminated water. In order to carry out the reticulation function of the pollutants, the CL, CS$_P$ and NP$_S$ structural components have been chemically functionalized with glutaraldehyde (F$_{CS}$) and epiyclorhidrine (F$_{CL}$), so that, we might obtain, by means of a compositing process, a material having a nanometric porous structure and usable in the molecular filtering processes (fig.2). The pollutants’ reticulation (F$_2$) at the level of the three structural components takes place in S$_2$ phase, each of them separating a certain category and type of target pollutants. Following the reticulation processes, it is likely that the reticulated pollutants should be partially or totally degraded, as a result of the chemical crosslinking interactions holding between the functional groups and the pollutants. The degradation function of the pollutants (D$_P$) is carried out at the level of all the structural elements of M$_{OF}$, and is due to the polyfunctional structure and to the nano- and micro- fiberlike morhosostructure of the composite. In order to degrade the pollutants (F$_3$), M$_{OF}$ was functionalized and composited, so that, due to the density of functional groups and due to the space distribution of its structural components, nano and microchannel structures may be formed, which may degrade and encapsulate the pollutants and the resulting degradation products, without offering the possibility for them to escape in the filtered environment. The controlled separation process of the depolluting products is done by means of the F$_4$ function, carried out by the NP$_S$ nanostructural component. NP$_S$ contains Fe$_3$O$_4$ nanoparticles functionalized with glutaraldehyde, which assures the separation of the composite filled with pollutants. These nanoparticles can be stimulated, in order to be separated in a controlled manner, by means of an external magnetic field (MS) (fig.2). The NP$_S$ structural component can reticulate organic pollutants that have chemical aminated functions, by means of the F$_{NP}$ functional structure.

![Diagram](https://www.intechopen.com)

**Fig. 3.** Typical structural and interfaceing diagram of a composite material (CM$_{OF}$, M$_{OF}$ – component material phase of M$_{OF}$; F$_1$, F$_2$ – functional elements of CM$_{OF}$; SF$_1$, SF$_2$ – microsurfaces distributions of functionalities; SI$_{C1-C2}$ – compositing interfaces of CM$_{OF}$)

The biodegradation function (F$_5$) is the result of the appropriate modelling of the morphological and morphochemical structure of M$_{OF}$, so that, when the composite charged with pollutants is exposed to the environment factors, it might be degraded and assimilated, without modifying the structure and the functionality of the environment. The CL structural component encapsulates the reticulated pollutants and the degradation products, so that the environment may assimilate more easily the M$_{OF}$ charged with pollutants, without generating short or long term toxic effects.
There is not a well defined limit between the intelligent materials and the functionalized materials, as in both cases material structures which are “programmed” to execute pre-settled tasks are involved. The polyvalent functionalization of $M_{OF}$ involves the discretization of the obtaining processes, the carrier material phases being integrated in stages and by different processes. The most sensitive stage in the obtaining process of $M_{OF}$ involves the preserving and quantification of the functional structure and of the carried out functions, so that they are not lost, masked or modified, following the compositing processes. The preservation and quantification of the functionality need the interposition of material phase connection, which should link the material structures that enter the composites structure ($CM_{OF}$) (fig.3). These connecting structures are called interface structures (IS) and have the role to protect and connect each structural component, as well as to preserve their functions. IS (fig.1) establish transition microregions between structural components (fig.3), the most widespread being the chemical interfaces.

![Fig. 4. General mechanism for functional structure masking of polyfunctionalized $M_{OF}$](image)

Chemical interfaces favour the interposition between two homogenous material phases, different in point of structure and physicochemical properties of the functional groups, coated layers or molecular phases. As it is difficult to find the structural interface which may realise, at the same time, the connecting and preserving functions of the functionality, in most cases, between the nanomaterial composites and its structural components, molecular layers interpose to meet one of the two imposed conditions. Depending on the application and the type of functionalised composite, IS has an important role in stabilizing the morphochemical and morphostructural functionality by establishing complex physicochemical equilibria, which influence the dielectric structure, chemical and thermical stability, electric conductivity, magnetic moment, radiative stimulation, molecular diffusion, etc. The dimensions, the morphological structure, the morphochemical structure and the manner of distribution of IS are strictly conditioned by the chemical structure of the interfaced material phases and by the functionality we seek to obtain. For instance, in the case of the magnetically stimulable nanostructures, the thickness of the coating layer may affect, by screening, the magnetical separation processes (fig.4). In order to obtain structural interfaces that would not mask or cancel the functionality of the composite, each material structure of the composite is obtained and functionalised separately, following that the compositing processes of the different structural components would be carried out in a controlled manner and in distinct stages. The functionalised nanocomposite materials have
become indispensable tools in most scientific and technological fields, due to the flexibility of the structural and functional modelling mechanisms, as well as due to the available manipulation and control possibilities. Environment science and protection (water depollution, waste control dispersion, water purification, water filtration, etc.), biochemical sensing, detection and nanomedicine are the most promising fields in which this type of materials can be applied.

2. Mono- and polyvalently functionalized nanocomposite materials

By definition, mono- or polyvalently functionalised nanocomposite materials (CM) are the carriers of one or more functionalities. The structural and functional modelling of composites depends on the type of application and functions that the composite must carry out. In depolluting applications, the polyvalently functionalised composites have the advantage that they may be structurally and functionally modelled in an oriented manner, so that they might reticulate, encapsulate and separate a large number of organic pollutants.

2.1 Functional structure modeling of orientated functionalized materials

Chemical functionalization compounds are those chemical compounds able to establish stable polydirectional chemical bonds, not only with the carrier support, but also with the target molecular structure. Some applications (analytical applications, molecular separations, etc.) constrain the use of $\text{MO}_x$, imposing that the reticulation processes should not degrade the crosslinked structure. Compounds with high chemical aggressiveness are not proper to be used in the analytical processes of sensitization, reticulation, or macromolecular separation, which are specific to the biological compounds, organic pollutants, nanopores, etc. Highly aggressive reticulation chemical compounds can degrade, alter and disperse the morphochemical structure of reticulated target structures.

The most used classes of chemical functionalization compounds are triazines (fig. 5), sulphochlorides (fig 6), periodates (fig. 7), cyanate compounds and their derivates (BrCN, ClCN, hydrazine - fig. 8), aldehydes (glutaraldehyde, N-hydroxysuccinimide, benzoquinone, etc. – fig. 9) and the epoxidic compounds (epichlorhydrine - fig. 10).

The glutaraldehyde is the most proper compound to be used in macromolecular functionalizing processes, as it can easily form stable functional structures, which do not degrade reticulated structures. The rest of the functionalization chemical compounds mentioned in the literature (antibodies, biological markers, functionalization compounds for the noble metals or for those with low reactivity, etc.) is characterized by the fact that they are specific to a certain carrier support layer and reticulate a strictly limited spectre of target structures. As functionalizable carrier supports, different types of polymeric nanostructures may be used (cellulose, dextrans, elastomers, glicols, amines, acetamides, etc.), nanostructured elements (oxidic structures, metals, fulerenes, quantum dots, etc.), other nanostructured materials having their surface and depth structure enriched with chemical groups (-H, -OH, -SH, $\text{H}_2\text{O}$, etc.) that may establish chemical reticulation reactions. The composing process of the structural components of composites may take place in one or more stages, depending on the functionality and the assigned functions, in relation to the affinity and chemical stability of each functional component, as well as depending on the morphological and morphochemical structure of the structural components and of the composite we seek to obtain.
Fig. 5. General mechanisms of the macromolecular reticulation of triazinic-functionalized carrier supports (hidroxy-enriched carrier support)

Fig. 6. General mechanisms of the macromolecular reticulation of halidic-functionalized carrier supports (carboxy-enriched carrier support)

Fig. 7. General mechanisms of the macromolecular reticulation of IO-4-functionalized carrier supports (poliol-enriched carrier support)

Fig. 8. General mechanisms of the macromolecular reticulation of cyan-functionalized carrier supports (hydroxy-enriched carrier support)

Fig. 9. General mechanisms of the macromolecular reticulation of aldehydic-functionalized carrier supports (amino-enriched carrier support)
2.2 Morphological and morphochemical structure of nanostructured materials

The surface and depth morphological and morphochemical structure of the orientedly functionalized composite materials is radically modelled by the reticulation and self reticulation processes, and by the local physical and chemical processes, established between the structural elements of the composite (self reticulation, induced mechanical tensions, magnetic and electric dipole moments, hydrogen bonds, molecular or atomic reticulation sites) (fig. 11).

The agglomeration processes are specific both for the nanostructured materials, and for the functionalized nanocomposite materials, due to self reticulation processes, surface impurities, the electrostatic and micromagnetic polarization, superficial tensions between solvents and the dispersed functionalized material. Self reticulation is due to functionalization imperfections, which lead to the formation of ununiformly functionalized surfaces that favour the occurrence of atomic and molecular sites, able to initiate and establish reticulation bonds with the chemical functions of the neighbouring morphostructures. $M_{OF}$ morphology is modelled by the morphology of its structural
component, by the compositing techniques and conditions, by the expression mechanisms of the functionality and by the molecular association processes at a nanotopological level. The functionality and the morphostructure of the structural components are conditioned by the physical and chemical equilibria established between the structural elements of the composite. The morphological structure of M_{OF} may be modelled by choosing the adequate compositing methods (reverse micelle, photolysis, hydrolisis, etc.) and by the strict control of conditions and of the compositing phases (temperature, solvation effects, rate of agitation, etc.). The morphological and morphochemical depth structure of the composites cannot be modified during the compositing process, and the external modelling factors may lead to loosing of the functionality of the composite or of its strutural components. Reverse micelle method is the most convenient technique of obtaining nanostructured materials and functionalized composites.

3. Nanostructured material interfaces

Material interfaces (MI) are material structures that favour the stable binding of two or more material structures having different chemical and physical properties. The surface chemical structure of most of the inorganic structures does not bear direct chemical functionalization processes, and the growth of material interfaces is needed (functionalization molecule layers, coating layers) which should favour stable chemical bonds between the involved material structures (fig.12). Impurities, microsurface defects and the reduced reactivity of microsurfaces are the main elements that condition the functionality of this type of nanostructures. In order to favour the physical and chemical interaction processes between the organic and inorganic material phases, in most cases, a series of processes and chemical compounds are used to modify their surfaces energetically and chemically: silans, acid or basic chemical treatments, chemical passivations, etc.

In the case of high functionalized oriented structure composites, each structural component of the composite needs the deposition of one interfaceable chemical structures having a certain specificity in relation to the rest of the compositated structures. Preliminary chemical treatments due to eliminate impurities and to modify the treated microsurfaces energetically aim to favour chemical reticulation reactions and to obtain uniformly distributed functional interfaces. The elimination of surface impurities is indicated in the case of oxidic nanomaterials and amorphous material structures, whose surface is likely to be impurified with traces of water, hydroxyl groups, metals, ions, etc. The most efficient method to eliminate these impurities from the surface of oxidic structures or of amorphous material structures consists of treating the interest surfaces with acid mixtures (HNO_{3}, H_{2}SO_{4}, H_{3}PO_{4}) (Lucas et al., 2007). As a result of the acid treatments, part of the treated support structure is partially degraded, and the metals and metal ions are removed. After having carried out the acid treatment, the composite nanoparticles are washed with solvents, which should remove the resulted degradation compounds. In most cases, acid treatments are followed by chemical treatments intended to passivate the modified nanostructures electrostatically. Organometallic salts are usually used (citrates, oxalates, etc.) in order to eliminate the electric charge density induced in the surface structure (Campos et al., 2002).

Prevention of the agglomeration and flocculation processes can be made by introducing some additional repulsion forces, so that the support phase, chemically and energetically modified, should be positioned in a chemical, electrostatic and micromagnetostatic equilibria with the host fluid. The spreading of the ionic phases in aquaeous host medium is influenced by the
presence of the residual ions ($\text{Na}^+$, ($\text{CH}_3$)$\text{NH}_3^+$, $\text{NH}_4^+$, $\text{NO}_3^-$, $\text{ClO}_4^-$, etc.) and by the dielectrical intrinsic structure of the crystalline structure of the dispersed phase, establishing electrostatic Van-der-Waals bonds with the amphoteric hydroxyl groups (Campos et al., 2002).

The favouring of dispersion may be done by inducing some negative (dispersion in acid environments) or positive (dispersion in alkaline medium) electric charges at the level of dispersed phase surface (Massart et al., 1995). The dispersion of the inorganic phase in organic host medium (cyclic hydrocarbons, paraffine oils, linear caten hydrocarbons, etc.) involves coating them with an amphiphilic molecule layer, in order to initiate the steric repulsion processes. Molecular reticulation prevention is due to the thermic agitation movement of the terminal hydrocarbons that lead to the formation of specific crystalline structures. The most used surfactants are sodium bis (2-ethylhexyl) sulphosuccinate (AOT), cetyl trimethyl ammonium bromide (CTAB), lipids (lecithin, phosphatidyl glycerol dihexadecyl phosphate, etc.), oleic acid, stearic acid, etc.

Silanes and cyanides are the most indicated chemical agents to modify the nanostructured surfaces chemically and energetically, so that they should establish easily stable chemical bonds with OH-, H-, -O-, M-O and M (M – metal atom) groups found on most of the material structure surfaces. Cyanates compounds (BrCN, CICN) favorise the forming of a large spectrum of reactive chemical groups on oxidic surfaces, of the type: nitrile: (-CN), isonitrilic, cianamidic (-$\text{(R)}_2\text{N-CN}$), azinic (C=N)-, isocianic (-N=C=O), carbodiimidic (N=C=N), aminated (primary, secondary, tertiary), compounds having nitrogen and mixed functionality. This type of reactive chemical groups have reticulation properties which are similar to those mentioned in figures 5-10. The organofunctional silanes reaction with –OH groups surfaces is a substitution reaction at the silicium atom, resulting silylated surfaces in which the Si atoms fix on the surface by covalent chemical bonds, via oxygen (fig. 13). These bonds may be established directly or in the presence of water, through of an intermediary silanol. The reactivity of the hydroxilated surfaces with the silanes decreases in the following order: $\text{Si-NR}_2 > \text{Si-Cl} > \text{Si-NH-Si} > \text{Si-O} = \text{CCH}_3 > \text{Si-OCH}_3 > \text{Si-OCH}_2\text{CH}_3$. Si-O-microsurface bonds are the critical factor that sustains the reaction, not only in aprotic conditions, but also in non aprotic ones.
Fig. 13. General structure of an organosilane organic-inorganic interface (1 – hidrolisable functionalities: -OH, NH2, O-O, etc.; 2 – space linkers; 3 - inorganic modified substrate)

Methoxysilanes are able to sustain reactions with the surfaces from an energetic and chemical modification point of view, not only in aprotic and non aprotic conditions, having lower reactivity compared to the ethoxylans and needing catalytic conditions for sustaining. An important control factor of the physical and chemical characteristics of the chemically modified microsurfaces is given by the length of the chemical linkers interposed between the organic functionalities and the Si atom fixed on the modified surface, imposing a series of specific chemical reactivity constrains (Jall et al., 2004; Chechik et al., 2000). The organic surface imposes a series of steric limitations on the accessibility of the functional groups which are placed in their vicinity, and the space linkers with appreciable length confer an increased mobility in order to functionalize the entire target surface. Aluminium and titanium form stable surface oxides by hydrolicitic processes. Their oxidized surfaces have enough -OH groups to allow the coupling of the organofunctional silanes in the same conditions with those applied to the silicides layers.

Copper, zinc and iron form surface oxides, which are instable from a mechanic and hydrolicitic point of view. The reticulation process of the silanes on these types of surfaces involves two or more silans, one of which is a chelate agent (polyamines, polycarboxylic acids). The second silane is chosen depending on the chemical reactivity of the organic component and on its capacity to initiate condensing reactions with the first silan. In the case of weak reactive metals (Ni, Au, Pt, etc.), the reticulation of silanes on their surface involves the formation of coordinative bonds (phosphines, mercapto, amino-functionalized silanes). In order to do this, a second silan must be used, which may present reactivity towards the organic component. Titan, zirconium and nickel form stable hydrides. The ability of several metals to form amorphous alloys with hydrogen is exploited in order to adsorb and coordinate the silicon hydrides on the metal surface. The functionalization hydride of silanes and the treated surface of the metal release hydrogen in the presence of bases or noble metals.

4. Physical and chemical properties of functionalized nanosurfaces

Nanocomposites and nanostructured functionalised materials are those materials which carry their own chemical and physical properties, showing them in their next vicinity. The physical and chemical characteristics of structural components and of their composites
depend completely on how they have been obtained: precursors, solvents, physical and chemical properties of the structural interfaces, temperature, pH, etc. The main processes responsible for the morphostructural, morphochemical and morphofunctional modification of depth material interfaces are: a. chemical or electrostatic adsorption sites of: neighbouring molecular radicals, metallic ions, molecules with high diffusion degree and small molecular mass (fig.14); b. hydrogen bonds or other similar chemical bonds (fig.14); c. the type, density and length of molecular space-linkers (fig.14); d. the type of chemical bonds established by the material interfaces with the organic and/or inorganic interfaced structures.

In the case of organosilanic interfaces, the chemical and electrostatic sites are due to functional groups of silane which carry out functions in the reticulation of the target structures and in the molecular or atomic sites which possess electric dipole moment. The structural modelling process is a dynamic one, taking place in the phase of nucleation of the organosilane sites and at the level of the growing surface of the laid-down functionalization interface. As a direct consequence, micro- and nanostructures empty sites (space) are generated in the depth structure, favouring the diffusion of charged ions and of the structures with lower molecular mass (fig.14). The impurities reticulated in the depth structure come from the reaction medium, as most silanization processes are catalyzed processes (alcohols, NH₄OH, etc.), that may generate in the reaction medium different molecular structures (RₙSi(OH)₄⁻ₙ, N⁺Hₐ(ΟH)ₘ (j+n+m=5), H⁺, -OH, H₃O⁺, etc.) that may diffuse and reticulate in the depth of the laid-down functionalized interface. Self assembled monolayer (SAM) material interfaces have the advantage that they suppress morphochemical and morphostructural processes, but they do not present the same variety and functional affinity by comparison with layered material interfaces. The obtaining of layered material interfaces with varied and stable functionality may be an advantage for a series of applications in detoxification, depollution and decontamination, as they may reticulate, encapsulate and host a wide spectrum of toxic compounds in their structure.

Interfaces with high content of hydrogen bonds and chemical bonds with hydrophobic character can reticulate and host large amounts of aminated, carboxylated, hydroxylated, hydration water, active hydrogen sites and molecular radicals. Using silanes of the NH₄Rₙ-Si(-O)- (n+m = 4) type in processes of deposition of functionalized interfaces leads to the formation of polymeric silanic structures of the n[SiO₁.₅₋R(NH₂)](NH₂)ₙδ type, enriched with -OH, amino, free water and reticulated H₃O⁺ functional forms (y characterizes the degree of cohydrolisis of Si(-O)-, δ characterizes the amount of aminated groups) (Iordache et al., 2009). The thickness of the laid-down organosilane layer and the amount of reticulated aminated groups depends on the type of the used silane, the cohydrolisis time and the type of the used catalyst. In the case of F₃O₄ nanoparticles coated with organosilane polymer by cohydrolisis of the (3-aminopropyl)trimetoxsilane, functionalized interfaces of the F₃O₄- n[SiO₁.₅₋(CH₃)₃(NH₂)](NH₂)ₙδ type have been obtained. On the basis of the analysis of the acquired experimental data by energy-dispersive X-ray spectroscopy (tab.1), the growth rate of the coated layer has been determined, as being modelled by the analytic relation (1). The graph representation of the analytic relation (1) is presented in figure 16.

Nano and micro-sites established in the depth of the laid-down organosilane coated layer favour the diffusion of the lower mass molecular compounds, due to the type of space-linkers and to the flexibility of Si-O-Si (120°<θ<180°) bonds (Tolstoy et al., 2003). This fact is due to the unhydrolizable character of molecular space-linkers, which orientate the neighbouring hydrolizable structures depending on the electric dipole moment, the
chemical interaction potential and the mechanic tensions established between them and their neighbouring macromolecules.

Fig. 14. General physical and chemical mechanisms who model the morphochemical and morphological structure of functionalized nanosurfaces (1 – positively charged molecular radicals; 2 – hydrogen bonds; 3 – free space sites filed with positive charged molecular radicals; 4 – access pathway for small molecules; 5 – positively charged nanosurfaces; 6 – molecules with strong electric dipole; 7 – negatively charged molecular radicals; 8 – positively charged topological nanosites; 9 – free space for negatively charged molecular radical retaining; 10 – bonds susceptible for breaking; 11 – inorganic interfaced nanostructure; 12 - space linkers; 13 – organic functionality; 14 - space linkers)

The size of the free micro-sites established in the depth of the coated layer has the magnitude of dimensions comparable with that of molecular space-linkers, modelling the mechanic tensions and the charge distribution. Functional organosilane groups (fig.14) could be inserted on the surface and in the depth of interfaced structures, using as silanization agent: silanols, amino silanes, siloxanes, epoxy silanes, carbinolic-functionalized silanes, acrylate/metacylate-functionalized silanes, carboxy silanes, mercapto silanes or anhydrides-functionalized silanes (Denmark & Sweis, 2002). Functional organosilanic groups may reticulate nucleation macromolecular structures of the n[SiO1.5-DŽ-RX]Y_{n6} type (X functional silanic group: -NH₂, -SH, OH, -COOH, etc.) or the sites or molecular radicals (Y) in the reaction volume, modifying the morphochemical and morphofunctional distribution, as well as the electric charge density established on surfaces of the deep micro- or nanosites of material interfaces. In the case of the aminosilanes catalyzed by NH₄OH, the following bonds Si-O-Si, Si-NH-Si, Si-O(NH³⁺)-Si, SiNH³⁺, [M(NH₃)ₙ]ᵐ⁺, result during the process of the n[SiO1.5-R(NH₃)](NH₃)ₙ₆ laid-down coated layer. These bonds can be easily broken by the molecular radicals present in the reaction medium, resulting new types of functional group: Si-OH, Si-NH₂, etc.
Crt.  | N (Wt%)  | O (Wt%)  | Si (Wt%) | Fe (Wt%) |
-----|----------|----------|----------|----------|
P1   | 0.00     | 60.45    | 17.62    | 21.93    |
P2   | 1.97     | 42.10    | 22.02    | 33.91    |
P3   | 3.93     | 60.79    | 8.56     | 26.72    |
P4   | 2.49     | 51.40    | 19.08    | 24.20    |
P5   | 2.90     | 53.09    | 26.98    | 13.02    |
P6   | 4.06     | 53.22    | 14.05    | 17.74    |
P7   | 1.04     | 55.87    | 18.47    | 22.30    |
P8   | 3.50     | 44.06    | 20.93    | 13.76    |

Table 1. Contents of Fe, Si, N and O found on SEM investigated microsurfaces

\[
\ln([\text{N}]+[\text{Fe}])t - \ln([\text{Si}][\text{Fe}]) = f(t) \quad (1)
\]

Fig. 15. \(\text{Fe}_3\text{O}_4\cdot n[\text{SiO}_{1.5}\cdot \text{(CH}_2\cdot 3(\text{NH}_2)](\text{NH}_2)n_6\) nanoparticles
(a) TEM image (180.000x) (b) SEM image

In the case of \(\text{Fe}_3\text{O}_4\) nanoparticles coated with \(n[\text{SiO}_{1.5}\cdot \text{(CH}_2\cdot 3(\text{NH}_2)](\text{NH}_2)n_6\), the experimental data we have obtained by electronic transmission microscopy (ETM), electronic scanning microscopy (ESM) and EDX prove that the morphostructural and morphochemical modelling takes place at the level of the laid-down organosilane layer and at the level of the \(n[\text{SiO}_{1.5}\cdot \text{(CH}_2\cdot 3(\text{NH}_2)](\text{NH}_2)n_6\) nucleation sites. This fact entails that chemical stabilization and passivation of the laid-down interfaces take place directly during the coating process, by microlocal passivation of nucleation sites and at the level of the growth surface, in the presence of the reaction medium content.

For biochemical reticulation applications, unstable chemical reticulated impurities on the surface and in the depth of coating layers can be removed by washing them with \(\text{H}_2\text{O}, \text{NaCl}\) or organomethalic salts (citrates), having the role to stabilize the suspension nanoparticles from a chemical and isoelectrical point of view. The obtained suspensions (4%) are used with the pH adjusted to 7±8 by proper dilution.
Analytical modelling of organosilane layer growth laid-down depending on cohydrolysis time

Experimental data fit: $y_0 + Ae^{x/t}$

Fig. 16. $f(t)$ function graph

**5. External stimulable nanocomposite materials**

Externally stimulable nanocomposites represent one of the latest priorities and needs in nanoscience and nanotechnology field, having applications in various fields, such as: chemical and biochemical detection (sensors, differentiated and undifferentiated prelevation of biochemical analytes, etc.) (Festag et al., 2005), nanomedicine (Murcia & Naumann; Bekyarova Haddon & Parpura; Gao; Zheng & Huang; Watanabe & al.; Meziani & al.; Lévy & Doty; Hattory & Maitani; Lellouche; Kommareddy, Shenoy & Amiji; 2005), environment science and protection (depollution, decontamination), chemistry (molecular and macromolecular separation). The stimulation concept designates that way by which a material structure registers a change in its energetic or structural structure under the influence of an external stimulation factor. In this sens, electromagnetic stimulation processes are very well know (fluorescence, phosphorescence, photocatalysis, semiconductor stimulation), magnetic (RMN, RES), radiative (elementary particles, nuclear and molecular radiation). By implementation of some control functions at the level of a stimulable material nanostructure, oriented controlled stimulable nanostructures can be obtained in order to measure, quantify and control the stimulable induced states. Polyvalently functionalized nanocomposite structures are hard to obtain, especially in reticulation applications with targets coming from systems with complex morphochemical and morphostructural, biochemical and biological organisation. Kinematic and dynamic control methods of the stimulable material structures placed in carrier host medium with complex molecular structure are relatively limited and reduced to control by magnetic, electric and electromagnetic stimulation. These nanoparticles have been functionalized so that they should reticulate undifferentiatedly and undestructively the target biological structures, which vary from a morphotostructural and morphofunctional point of view.
Analytical techniques with constant metrics use only a stimulation factor for the sensitization of a large spectrum of analytes. Existent analytical methods for biological sensitization (Bhunia, 2008; Jasson, 2010) are variant in respect of a specific required stimulation factor, being unfriendly in respect of sensitization of a large analyte spectrum.

The analytical method proposed in figure 17 starts from the assumption that the set of values formed by the magnetic moment attached to biochemical structures (m) by reticulation processes and the mass (M) attached to reticulation macrocomplexes are uniquely determined for each type of biochemical structure. As the prelevation and magnetic discrimination processes of the analytes take place in a host carrier fluid, we started from the assumption that the friction force \( F_\mu \), weight force \( F_G \), archimedic force \( F_A \) and micromagnetical force \( F_m \) (fig.17) are the only ones that can model the kinematics and the dynamics of the magnetic discrimination. In order to reduce the number of the observables and degrees of freedom that affect the kinematics and the dynamics of the \( \text{Fe}_3\text{O}_4 \) nanoparticles reticulated on the surface of the targeted biostructures (NPB), a force field configuration was proposed (fig.17a, fig. 17b), in which NPB movement is orientated, as follows: a. parallel with the gravitational acceleration vector (fig.17.b); b. opposed to the gravitational acceleration vector (fig.17.a). The observables attached to the configured biochemical sensitization process are the specific speeds \( v_i \) of NPB movement in magnetic field and the specific surface fluorescence of the reticulated biochemical structures \( \tau_i \), so that an univoque analytical relation can be established between the discriminated biostructure and its specific surface macromolecular fluorescence mark. From an analytical point of view, the set of observables \( (v_i, \tau_i) \) can be correlated by software, so that, for each \( v_{i1} \) (a), \( v_{i2} \) (b) of sensitized NPBs, according to the configurations proposed in figure 17, one may identify those sequences that present the same fluorescence mark \( \tau_i \), according to relations (4) and (5). The index „i” designates the type of the sensitized biochemical structure and is defined on the entire reticulation and sensitivization-detection spectrum. Analytical equations (4) and (5) model and describe the proposed constant metric analytical model, which is deducted from fundamental equations (2) and (3), which describe the NPB movement in configurations 17a and 17b.

\[
F_{ARH} + F_F = F_{EM-1} + Mg
\]
\[ \text{F}_{\text{EM-2}} + \text{F}_{\text{ARH}} = \text{Mg} + \text{F}_F \]  

where: \( \text{F}_{\text{ARH}} \) is the archimedic force, \( \text{F}_{\text{EM-1/2}} \) are the magnetic field gradients, \( \text{F}_F \) is the friction force opposed to NPB movement in the host fluid, \( \text{F}_F = -\text{kx} \), \( \text{M} \) is the mass of NPB.

\[ M_{\text{NPB}}(i) = 2m\left(\frac{\text{dB}_1}{\text{dx}_1} - \frac{\text{dB}_2}{\text{dx}_2}\right) / \sqrt{2c^2v_f^5 \left( \frac{1}{x_2} - \frac{1}{2x_1} \right)} \]  

\[ M_{\text{NPB}}(i) = m\left(\frac{\text{dB}_1}{\text{dx}_1} - \frac{\text{dB}_2}{\text{dx}_2}\right) / \left\{ \frac{\sqrt{2v_f^5v_1}}{2x_2} + \frac{\sqrt{2v_f^3v_2}}{2x_1} \ln \frac{v_1 + (v_2 - v_1)e}{v_2} \right\} - \frac{2v_f}{4x_1} \]  

where: \( \text{dB}_1/\text{dx}_1 \) and \( \text{dB}_2/\text{dx}_2 \) are the gradients of the magnetic field realized on the sensitization distance, \( x_1 \) and \( x_2 \) are the sensitization distances, \( v_1 \) and \( v_2 \) are the characteristic speeds of each NPB, \( v_1 \) is the speed of the carrier fluid which hosts the NPB.

In the above mentioned conditions, equation (4) describes those NPB that, following the magnetic acceleration process, achieve a constant speed (c), at the moment of the sensitization of discrimination and for a certain value of the field gradient (dB/dx). Equation (5) is specific for those NPB that, after the magnetic acceleration process, did not achieve a constant limit speed (c). According to equations (4) and (5) it results that analytical methods with invariant metric may be configured for biological detection and monitoring by selective magnetic discrimination. Also, according to these equations, it results that the sensitization domain may be improved by modifying the magnetic field parameters (dB/dx) of the external stimulation factors. The magnetic field represents a convenient method for the undestructive stimulation of magnetic dipole carrier functionalized material structures, which, having a considerable action range, as compared to the rest of the known stimulation factors. Due to the fact that most organic compounds are diamagnetic, the magnetic field substantiates new undestructive selective techniques for controlled stimulation, as the stimulable material nanostructures are not masked by their neighbouring material structures.

6. Structure and functionality modelling of nanocomposite materials designed for undestructive reticulation of biological structures

The functionalization of nanoparticle and nanocomposite structures for biochemical reticulation requires a set of conditions involving: the specificity and the recognition of the reticulated structure, the chemical and functional stability, avoidance of the biochemical interaction with the host carrier, avoidance of self reticulation and agglomeration processes. Most of the applications, especially the analytical one, require undestructive biological and biochemical reticulation in respect to the targeted biochemical structure. Glutaraldehyde is the chemical functionalization and reticulation compound frequently used in the analytical biological process (fig.10), as it presents a weighted chemical aggressiveness and it preserves the internal morphochemical struture of the reticulated structures. Glutaraldehyde is a dialdehyde whose functional molecular residue (CHO) can reticulate undifferentiatedly the hydrogen active, amino- and tiol- sites, present on the surface and in the depth structure of most of biochemical structures. Polyvalent functionalizations require a series of cautions intended for the chemical stabilization of the functionalized nanostructures, so that different
implemented functional groups may not mask or cancel each other’s functionality and functions. In order to investigate this aspect, Fe$_3$O$_4$ nanoparticles have been coated with n[SiO$_{1.5\gamma}$-(CH$_2$)$_3$($\text{NH}_2$)]($\text{NH}_2$)$_{\text{ns}}$ and functionalized monovalently (with glutaraldehyde - GL) and polyvalently (with glutaraldehyde and epichlorhydrine - GL+ECH) (Jordache et al., 2009). For their morphochemical structure, the following structural relations have been proposed Fe$_3$O$_4$-[n[SiO$_{1.5\gamma}$-(CH$_2$)$_3$($\text{NH}_2$)]($\text{NH}_2$)$_{\text{ns}}$-(GL)$_{\text{ne}}$] (np-(GL)$_{\text{ne}}$), (ECH)$_{\text{nf}}$-[Fe$_3$O$_4$-[n[SiO$_{1.5\gamma}$-(CH$_2$)$_3$($\text{NH}_2$)]($\text{NH}_2$)$_{\text{ns}}$]+(GL)$_{\text{ne}}$ ((ECH)$_{\text{nf}}$-(np-(GL)$_{\text{ne}}$) (σ and β - modelling coefficients for functional fractions CHO, respectively Cl-C and -CH(O)CH -) (Jordache et al., 2009). For the functionalization of nanoparticles, 4 grams of chemically stabilized coated nanoparticles have been used (4% suspension), in the presence of 7 ml of glutaraldehyde (25%) ((6)). Polyvalently functionalized particles have been obtained by the functionalization of np-(GL)$_{\text{ne}}$ with an epichlorhydrine solution, previously prepared (NaOH, H$_2$O, acetone and 0.5 ml epichlorhydrine) (fig.9).

$$\text{Fe}_3\text{O}_4 - n\left[\text{SiO}_{1.5\gamma} - (\text{CH}_2)_3(\text{NH}_2)\right](\text{NH}_2)_{\text{ns}} + n\text{C}_5\text{H}_8\text{O}_2(\text{GL}) \rightarrow \text{NaOH} \quad \text{Fe}_3\text{O}_4 - n\left[\text{SiO}_{1.5\gamma} - (\text{CH}_2)_3(\text{NH}_2)\right](\text{NH}_2)_{\text{ns}} -(\text{GL})_{\text{ne}}$$

Glutaraldehyde functionalized nanoparticles have been exposed to air in order to investigate the undifferentiated reticulation potential of the saprophyte microorganisms, as the prelevated samples have been investigated by TEM (fig. 18). The same type of investigations has been carried out on B. Cereus, St. Aureus, E. Coli, Ps Aeruginosa, and the obtained results have been presented in figure 19. Also, the biochemical reticulation potential of np-(GL)$_{\text{ne}}$, and (ECH)$_{\text{nf}}$-(np-(GL)$_{\text{ne}}$ has been investigated, by ricin reticulation (Petrea et al., 2009). The samples have been investigated by scanning electron microscopy (fig. 20) and by confocal microscopy (fig.23). Before the investigations were performed by confocal microscopy, the functionalized nanoparticles were marked fluorochromically with rhodamine B, right in the phase of the laying down of the coated layer (cyan color in fig.23). The ricin was marked fluorochromically with fluorescein isothiocyanate (FITC) (green colour in fig.23). The results of the biological and biochemical reticulation investigations have proved that the obtained functionalized nanoparticles present a well defined functionality, and that the morphochemical structure of the reticulated structures has not been degraded (fig.18, fig.19).

![Fig. 18. Investigation images of the np-(GL)$_{\text{ne}}$ biochemical reticulation potential on saprophyte microorganisms (images acquired by Philips S208 TEM microscope)](www.intechopen.com)
It was observed that np-(GL)$_{n_e}$ are specifically distributed on the reticulated surfaces, according to the surface morphochemical structure of microorganisms. Most likely, distribution maps of the reticulated nanoparticles follow the patterns of the morphochemical distribution of aminated, thiolated, carboxylated, and carbonilated functional groups placed on the surface of microorganisms. The obtained data emphasize the np-(GL)$_{n_e}$ potential in the field of specific biochemical reticulation, as nanoprobes, or in the field of undestructive biochemical investigation (fig.18).

Fig. 19. Investigation images of np-(GL)$_{n_e}$ biochemical reticulation potential on B. Cereus, St. Aureus, E. Coli, Ps. Aeruginosa (images acquired by ESEM XL30 SEM microscope)

Using the elementary morphological segmentation method (Iordache et al., 2009), in order to determine the elementary morphostructural parameters, it was proved that the main agglomeration mechanism is micromagnetic. Also, it was proved that the functional structure of the obtained functionalized nanoparticle suspension is stable, and it does not initiate self-reticulation bonds. In the case of np-(GL)$_{n_e}$ and (ECH)$_{n_f}$-np-(GL)$_{n_e}$, which include Fe$_3$O$_4$ nanoparticles obtained by coprecipitation, the agglomeration domains present a spherical form (fig.20a, fig.20d, fig.21b). In the case of functionalized particles containing Fe$_3$O$_4$ obtained by reverse micelle technique, the agglomeration domains present an acicular form (fig.20b, fig.20c, fig.21a). These results prove the polyvalent functional structure of...
ECH\textsubscript{n,\,np\,-\,(GL\textsubscript{ne}), according to the specific functionalization mechanisms proposed in figure 22b.

Fig. 20. Investigation images of (ECH\textsubscript{n,\,np\,-\,(GL\textsubscript{ne}), and np\,-\,(GL\textsubscript{ne}) biochemical reticulation potential on ricin (images acquired by VEGA II LMU SEM microscope)
(a) np\,-\,(GL\textsubscript{ne}) (Fe\textsubscript{3}O\textsubscript{4} obtained by coprecipitation) (b) np\,-\,(GL\textsubscript{ne}) (Fe\textsubscript{3}O\textsubscript{4} obtained by reverse micelle) (c) (ECH\textsubscript{n,\,np\,-\,(GL\textsubscript{ne}) (Fe\textsubscript{3}O\textsubscript{4} obtained by reverse micelle) (d) (ECH\textsubscript{n,\,np\,-\,(GL\textsubscript{ne}) (Fe\textsubscript{3}O\textsubscript{4} obtained by coprecipitation)
Reticulation investigations carried out on ricin have proved that there is no significant difference between the reticulation mechanisms of microorganisms and those of ricin. Ricin reticulation tests have proved that the reticulation processes specific to toxins are degenerated, and a single functionalized nanoparticle crosslinks more macromolecules on ricin by contiguity.

The results demonstrate that the morphological and morphochemical structure of macromolecular biological structures play an important role in the evolution of reticulation processes, conditioning the mechanisms and their specificity, as well as the morphological and morphochemical structure, physicochemical properties and the stability of the resulted agglomeration domains. Morphochemical structure relations and the proposed functionalization mechanisms (fig.22) have been deduced by taking into account the obtaining conditions of the structure functionalities and the experimental data acquired during the chemical and biochemical reticulation investigation processes. Chemical and functional stability of the functionalized nanoparticle suspension, as well as the uniformity and the specificity of the biochemical reticulation processes, can be explained only if we accept the fact that the layer of functionalizing molecules, laid down on the coated layer, is continuous and uniform.

![Diagram](image)

**Fig. 21.** The adopted elementary and bulk morphostructural domains for (a) (ECH)$_{n_1}$-np-(GL)$_{n_2}$ (Fe$_3$O$_4$→MI) and (b) (ECH)$_{n_1}$-np-(GL)$_{n_2}$ (Fe$_3$O$_4$→CP) reticulated nanoparticles on ricin surface (W$_b$, W$_l$, L$_t$, L$_l$ = geometrical dimension of elementary and bulk domains)

The discontinuities on the functional structure of surfaces involve the existence of more different biochemical reticulation sites, favouring self-reticulation processes of nanoparticles in suspension. The acquired experimental data disprove the existence of masking, neutralisation or repeal processes between the CHO, CCl or CH(O)CH functional groups. Most likely, a stable chemical equilibrium is established between the CHO, CCl and CH(O)CH functional groups, as they are distributed on the surface of functionalized nanoparticles, according to a well determined morphochemical distribution map.
Fig. 22. Proposed morphochemical and morphofunctional structure for (a) npa-(GL)$_{me}$ nanoparticles (a): 1 - GL; 2 - propyamine chemical residues; 3 - the most probable morphochemical distribution of chemical functionalities in the depth of coated layer; (b): 1 - CHO functionalities; 2 - Cl functionalities; 3 - epoxy functionalities; 4 - masked CHO functionalities)

7. Structure and functionality modelling of nanocomposites intended for ecological depollution

Research in the waste water and solid waste management usually encounters three main challenging problems: the huge quantities to be processed, the large varieties of contaminants and impurities, and the conversion of solid residuals coming from specific treatments into chemically and biologically inert materials for their safe disposal. Additionally, the useful compounds recycling are preferential design target for most of researches, but in many cases the safe disposal of inactivated residuals is prevailing. In spite of the many advances in waste water treatment, the problem of removal the organic pollutants is still pending around the elaborated technologies developed under the tertiary waste waters treatment concept. Thus, oxidative technologies are expensive and degradation products are still polluting (Gogate & Pandit, 2004). Flocculation and coagulation (Cheremisinoff, 2002) as well as ionic exchange technologies (Robinson & al., 2001), are limited in their efficiency for the removal of organic compounds. Chlorination is often use in the treatment of organic polluted waters, but its drawback is well known, when a large number of pollutants are converted into other compounds with the same toxicity. Microbiological technologies (Tedder & Pohland, 1990) and photocatalytic technologies (Hashimoto & al., 2005) are very selective ones and totally inappropriate for a broad spectrum of organic pollutants in the treated waste waters. Membrane materials (Cassano & al., 2001), particulate nanomaterials and functionalized hybrid nanomaterials as well as functionalized composite material adsorbents seems to be the best solutions for treatment of waste waters highly contaminated with a large number of organic compounds from different classes. Only the prices of these materials are restrictive for
their application at larger scale. The main factors that restrict depollution methods and technologies of surface waters charged with organic pollutants are due to: a. a great number (spectrum) of organic pollutants (OP) contained by contaminated waters; b. morphostructural and morphofunctional varieties of pollutants; c. dispersion mechanisms of pollutants in the environment; d. lack of controlled separation possibilities of pollutants from the source of pollution; e. lack of control upon degrading products, resulted during depollution processes; f. collecting, stocking, storage and management depollution waste.

Fig. 23. Investigation images of \((\text{ECH})_{n\theta}-\text{np-(GL)}_{n\varepsilon}\) and \(\text{np-(GL)}_{n\varepsilon}\) biochemical reticulation potential on ricin (confocal investigation using a multifoton broadband Leica TCS SP 2 microscope) (a) \(\text{np-(GL)}_{n\varepsilon}\) \((\text{Fe}_3\text{O}_4\) obtained by coprecipitation) (b) \((\text{ECH})_{n\theta}-\text{np-(GL)}_{n\varepsilon}\) \((\text{Fe}_3\text{O}_4\) obtained by coprecipitation) (c) \(\text{np-(GL)}_{n\varepsilon}\) \((\text{Fe}_3\text{O}_4\) obtained by reverse micelle) (d) \((\text{ECH})_{n\theta}-\text{npa-(GL)}_{n\varepsilon}\) \((\text{Fe}_3\text{O}_4\) obtained by reverse micelle)

The spectrum and the morphostructural and morphofunctional variety (SVM) specific to the content of organic pollutants are due to the variety and diversity of the materials used in industrial and domestic activities. SVM limitation, control and determination is a complex issue, which cannot be solved by conventional procedures, as it is conditioned by the exigence, necessities and material needs of the society. From the point of view of the morphochemical type and variety, surface contaminated waters contain inorganic pollutants (salts and metal derivates, mineral acid salts, mixed composition suspensions and amorphous material structures, simple metallic oxides, mixed metallic oxides, etc.), carrier vectors of radioactive elements, organic pollutants (hydrocarbons, alcohols, phenols, derivates of benzen, organic acids and their derivates, carboxylic compounds and their
derivates, amines, organic halogens and their derivates, aromatic derivates, organometallic compounds, detergents, etc.), biological pollutants (bacteria, viruses) and biochemical pollutants (toxines, proteins, peptines, sugar, etc.) (Evanghelou, 1998). Most of the organic pollutants degrade in time, generating other polluting compounds that may present high toxicity. Not only the organic pollutants, but also their degradation products, may affect on long and medium term human health and the functionality of the environment.

The dispersion and the diffusion of pollutants is constrained by their physical and chemical properties. Once introduced in the external water circuit, pollutants become uncontrollably, randomly and irreversibly dispersed on wide surfaces. Due to the complexity of the above mentioned factors, the separation processes of the organic pollutants in surface waters and industrial contaminated flux water will be carried out incrementally, using complex technologies and solutions. In most cases, separation processes are only realised partially, as used technologies and materials are oriented towards the removal of certain pollutants or of some limited classes of organic pollutants.

Oxidic materials with oriented functional structure (M_{OF}) offer promising development and new implementing solutions for the development of new methods to control depollution of contaminated waters, due to the modelling possibilities of functional structure, and to extend the spectrum of separate pollutants.

**Fig. 24. General mechanisms for the retention, degradation and encapsulation of pollutants**

1 - Su; 2 – functionalised magnetite; 3 - OH functional groups; 4 - amino-functionalities; 5 – epoxy-functionalities; 6 – fiberlike cellulose; 7- ways of pollutant degradation; 8 – retention of pollutants; 9 – paths for pollutant diffusion in the depth of composites; 10 – free space for encapsulation; 11 – pollutants; 12 – paths of pollutant host carrier in the depth of composite

M_{OF} separated pollutants spectrum depends directly on the type, polyvalency and stability of the obtained functional structure. Natural oxidic materials, with mixed nanostructures are the ideal candidates in order to obtain polyvalently functionalized materials, due to the complexity of the surface chemical structure, rich in reactive molecular fractions: -O-M, M-OH, H, -O-O-, hydrolizable molecular residue, electrically charged sites, embedded organic compounds, etc. Moreover, for the applications involving depollution, as well as for the
obtaining and functionalization of M_{OF}, there is no need for special chemical and technological conditions, as their usability is constrained by the enrichment and the stabilization of the functionality structure. The stability of this type of materials involves the morphochemical and morphofunctional stability, in order to preserve the morphostructural, morphochemical and morphofunctional characteristics, as well as to prevent the degradation of the support material, carrier for the reticulated pollutants.

In most depollution situations, the chemical and morphostructural stabilization of the functionalized composites is done under water, so that the depolluting applications are conditioned by the possibility of the composite to reticulate the pollutants in free chemical conditions (fig.5÷10). According to the above proposed morphostructural and morphofunctional modelling mechanisms, a polyvalently functionalized composite has been obtained. This composite integrates in its structure a mixture of natural oxides (Su), functionalized with BrCN (~100 grams, about 10 grams of np-(GL)_{n} nanoparticles and 100 grams of cellulose acetate functionalized with epichlorhidrine ((7)). The chemical composition of Su was determined by Wavelength Dispersive X-ray Fluorescence (WDXRF), and it was found that Su contains the following main fractions: SiO_{2} (70.13%), Al_{2}O_{3} (14.33%), Fe_{2}O_{3} (4.36%), K_{2}O (2.78%), Na_{2}O (0.46%), MgO (0.87%), P_{2}O_{5} (0.26%), TiO_{2} (0.58%), CaO (0.87%).

Fig. 25. Morphological and morphochemical structure of M_{OF-DP}
The composite presents a nano-fiberlike character (fig. 24, fig. 25a), due to the nano- and microstructural character of the water regenerated cellulose. Functionalized cellulose acetate performs the reticulation, compositing and spatial distribution functions of the Su and functionalized magnetite, as structural components. The structural elements of the composite are spatially distributed, on the surface and in the depth of cellulose microfibers, so that they lead to the formation of nano- and micro-membranary structures. Using the elementary segmentation method (fig. 25b), the mean distribution values of the elementary morphostructural parameters have been determined: perimeter (6.78 nm), length (17 ÷ 924 nm), area (3.85 nm²) (fig. 25d). The morphostructural character of the nanocomposite allows that the contaminated water may be filtered, at molecular level, from a mechanic, chemical, biochemical and biological point of view, according to the reticulation, degradation, and encapsulation mechanisms proposed in figure 24. It was noticed that the composite allows the development of the saprophyte bacteria on its surface (fig. 26), and points out the stable character of the material functionality and the biodegradability of its composite structure. Moreover, the biodegradable nature of the composite is also given by the nature of its composite structure, as its structural elements can be assimilated without inducing toxic effects in relation to the structure and functionality of the environment.

Fig. 26. Growth and develop of saprophite microorganisms on surface of M_{OF-DP}
functionalization processes of surface oxide components with BrCN. Most likely, in the first stage BrCN hydrolyses, leading to the formation of HCN, which attacks metal ions, anions and electric charged microsites.

Fig. 27. IR spectra of composite and their structural components
(a) IR spectra of: citrate passivated Fe₃O₄ nanoparticle (P1-Cit), Fe₅O₄·mSiO₁.₅(CH₂)₃(NH₂)₃(NH₂)₆(Gl)ₙ(P1-F), functionalised cellulose (P4), Su and MOF-DP (P3) (b) 1000-1550 cm⁻¹ IR spectra of composite (c) 500-800 cm⁻¹ IR spectra of composite

The CN⁻ anion interaction with surface oxides leads to the formation of a wide variety of functional groups, such as: NXₘYₙ→H (hydrogen bonds), N⁺Xₘ→OH⁻ⁿ (i = 0, 1, j = 1±3, n = 0÷2), water-hydrogen bonds), NH₃, CN, =C=NH, =C=NXₘYₙ, NXₘYₙ, C⁺ₘN⁺Xₙ (n+m=5; X, Y = H, -OH, halogens, other metallic ions). In return, aminated functionalities interact with metal ions in the structure of oxidic components, resulting new types of functional mixed structures, such as: (FeOFe, FBrO₂, OCIF₄) (506 cm⁻¹), (Fe-O₂) (503 cm⁻¹), peroxides (920-750 cm⁻¹), O=CCIF (501 cm⁻¹), O=SiCl₂ (501.1 cm⁻¹), C-Cl (501.9 cm⁻¹), (ClHBr, P₂O₆) (508 cm⁻¹), (FNCl, Al(OH)₄, CdO, CINNN, VF₅) (720 cm⁻¹), ((ORel₄)·(Si₂O₆) (508 cm⁻¹), (HCN)(721 cm⁻¹), (LiNC)(722.9 cm⁻¹) (Nakamoto, 2009, 2009). Most likely, the
shape of the absorption band 950-550 cm\(^{-1}\) is due to the significant contribution of the absorbing oxide forms, of the following types: MO\(^4\): (SiO\(^4\))\(^4\) (819, 596 cm\(^{-1}\)), (PO\(^4\))\(^3\) (938, 567 cm\(^{-1}\)), (SeO\(^4\))\(^2\) (822, 856 cm\(^{-1}\)), (ClO\(^4\))\(^-\) (928 cm\(^{-1}\)), (BrO\(^4\))\(^-\) (878, 801 cm\(^{-1}\)), (TiO\(^4\))\(^4\) (770, 761 cm\(^{-1}\)), (ZrO\(^4\))\(^4\) (846, 792 cm\(^{-1}\)), (HfO\(^4\))\(^4\) (800, 796 cm\(^{-1}\)), (VO\(^4\))\(^-\) (818, 780 cm\(^{-1}\)), (VO\(^4\))\(^3\) (826, 804 cm\(^{-1}\)), (CrO\(^4\))\(^2\) (863, 833 cm\(^{-1}\)), (CrO\(^4\))\(^3\) (844, 775.8 cm\(^{-1}\)), (MoO\(^4\))\(^2\) (897, 837 cm\(^{-1}\)), (WO\(^4\))\(^2\) (931, 838 cm\(^{-1}\)), (SiO\(^4\))\(^4\) (819, 596 cm\(^{-1}\)), (MnO\(^4\))\(^2\) (902, 834, 820, 821 cm\(^{-1}\)), (MnO\(^4\))\(^3\) (789, 778 cm\(^{-1}\)), (TcO\(^4\))\(^-\) (921 cm\(^{-1}\)), (ReO\(^4\))\(^-\) (971, 920 cm\(^{-1}\)), (FeO\(^4\))\(^2\) (832, 790 cm\(^{-1}\)), (RuO\(^4\))\(^2\) (845, 830 cm\(^{-1}\)), (RuO\(^4\))\(^2\) (840, 804 cm\(^{-1}\)), OsO\(_4\) (965.2, 960.1 cm\(^{-1}\)), (CoO\(_4\))\(^4\) (670, 633 cm\(^{-1}\)) (Nakamoto, 2009). The absorption peaks placed at 1645, 1710, 3335 cm\(^{-1}\), can be assigned to C = O functional groups, which are characteristic for polyurethane polymeric structures, resultated from the reticulation of -OH with aminated residues, placed on the surface of oxides. The absence of the absorption peaks of C=O, suggest that glutaraldehyde and functionalized Fe\(_3\)O\(_4\) nanoparticle were entirely reticulated by the functionalities containing nitrogen during material composition processes. Chemical reticulation processes have favoured the compositing process, by embedment of functionalized nanoparticles of magnetite and Su on surface and in the deep structure of cellulose.

8. Conclusions

This chapter presents the main physicochemical and functional mechanisms substantiating the understanding and proper modelling of the process of obtaining composite materials with functional oriented structure. These mechanisms were substantiated on the theoretical and experimental data obtained by the team of authors and their collaborators inside applicative projects for the sensitising and detection of biochemical structure and for the separation of organic pollutants in contaminated waters.

The chapter proposes a new analytical method with invariant metrics for sensitization and biochemical monitoring, based on using magnetically stimulable functionalized nanostructures, able to prelevate a great number of biochemical analytes, undifferentiatedly and undestructively. Biochemical invariance sensitization-detection is provided by imposing analytical and practical constraints at the level of analytical acquisition and processing systems, according to the configuration of the magnetic field of the discriminator, molecular weight, specific biochemical fluorescence spectrum. We have presented the analytical structure and the elements that model the process for obtaining magnetically stimulated functionalised composites, orientated towards the detection and undestructive biological reticulation.

The second part of this chapter presents the modelling and the obtaining of new polyvalently functionalized types of materials oriented to the reticulation, encapsulation and separation of organic pollutants in contaminated waters. The obtained material has a strong, biodegradable character, being modelled morphostructurally and morphofunctionally, so as to separate by reticulation a large number of organic pollutants, toxins and microorganisms. The chapter describes the main physicochemical mechanisms that model the conditions of separation, degradation and encapsulation of pollutants.

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