Method of controlling fluid loss in a permeable formation

Verfahren zur Kontrolle des Flüssigkeitsverlusts in einer durchlässigen Formation

Procédé pour contrôler la perte de fluide dans une formation perméable

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Description

[0001] The present invention relates to a method of controlling fluid loss in a permeable formation penetrated by a wellbore.

[0002] During the drilling of an oil well, a usually aqueous drilling fluid is injected into the well through the drill pipe and recirculated to the surface in the annular area between the wellbore wall and the drill string. The functions of the drilling fluid include lubrication of the drill bit, transportation of cuttings to the surface, overbalancing the formation pressure to prevent an influx of oil, gas or water into the well, maintenance of hole stability until casings can be set, suspension of solids when the fluid is not being circulated and minimizing fluid losses into and possible associated damage/instability to the formation through which drilling is taking place.

[0003] Proper overbalancing of formation pressure is obtained by establishing fluid density at the desired level usually via the addition of barite. Transportation of cuttings and their suspension when the fluid is not circulating is related to the fluid viscosity and thixotropy which depend on solids content and/or use of a polymer. Fluid loss control is obtained also by the use of clays and/or added polymers.

[0004] Fluid properties are constantly monitored during the drilling operations and tailored to accommodate the nature of the formation stratum being encountered at the time. When drilling reaches the producing formation special concern is exercised. Preferentially, low solids content fluids are used to minimize possible productivity loss by solids plugging. Proper fluid density for overbalancing formation pressure may be obtained by using high salt concentration aqueous brines while viscosity and fluid loss control generally are attempted by polymer addition.

[0005] When high permeability and/or poorly consolidated formations are penetrated as the zone of interest, a technique referred to as "under-reaming" often is employed in the drilling operations. In this process, the wellbore is drilled through the hydrocarbon bearing zone using conventional techniques and drilling muds. A casing generally is set in the wellbore to a point just above the hydrocarbon bearing zone. The hydrocarbon bearing zone then is redrilled using an expandable bit that increases the diameter of the hole. The purpose of the under-reaming is to remove damage from the permeable formation introduced during the initial drilling process by particles of the drilling mud and to increase the exposed surface area of the wellbore. Typically, under-reaming is effected utilizing special "clean" drilling fluids to minimize further formation damage. The high permeability of many hydrocarbon zones allows large quantities of the clean drilling fluid to be lost to the formation. Typical fluids utilized in under-reaming comprise aqueous brines which are viscosified with a polyacrylamide polymer to aid in the removal of the drill cuttings. US-A-3,818,998 discloses certain water-dispersible cellulose ethers, polysaccharides and polyacrylamides which may be crosslinked by contacting with a hexavalent chromium compound and a polyhydric alcohol to form a rubbery gel which swells in the presence of water. The gels may be used in the control of fluid loss. To control fluid loss, solids such as ground salt crystals, calcium carbonate or the like may be added to the fluid to form a filter cake on the face of the formation. This filter cake must be removed prior to or after the zone is gravel packed if hydrocarbon production is to be achieved. The salt crystals can be removed by circulating unsaturated salt brine to dissolve the particles. If this occurs before gravel packing, the circulating fluid often causes sloughing of the formation into the wellbore and yet further loss of fluids to the formation. If removal is attempted after the gravel pack, the gravel packing material often traps the particles against the formation and makes removal much more difficult. Other particulates, such as the carbonates can be removed with circulation of acid, however, the same problems may arise.

[0006] Completion fluids are those fluids used after drilling is complete and during the steps of completing the well. Completion can include cementing the casing in the well, perforating the well, setting the tubing, pump and the like. Completion fluids are generally used to control the well pressure, provide fluid loss control, prevent the collapse of tubing from overpressure and to reduce or prevent corrosion of the casing when left in a wellbore as a packer fluid above the production packer. In such an instance the fluid is placed in the annulus between the casing and production tubing string above the production packer to seal the annulus against the influx of formation fluids. Because a packer fluid may remain in a wellbore for many years before any remedial work is needed, such a fluid generally should be temperature stable, non-settling, non-corrosive and non-damaging to the formation.

[0007] Stimulation of hydrocarbon bearing formations typically is effected to increase the exposed area of a subterranean formation for flow of a hydrocarbon to a wellbore. Commonly, fracturing operations are performed to produce conductive channels from the wellbore into the formation. The channels may have a particulate proppant introduced therein to assist in maintaining the channels open within the formation. While creating fractures in low permeability formation is often considered a routine technique, the formation of fractures in higher permeability formations (500 millidarcy and greater) presents several problems. Effective fluid loss control is essential to creation of a fracture of any length in a high permeability formation. Typical fluid loss control additives for stimulation fluids comprise the ground salt, ground calcium carbonate and the like previously discussed in regard to clean drilling fluids. These materials are difficult to remove from the created fractures, particularly after the fracture is propped open by the introduction of a propping agent. The presence of unremoved fluid loss additives can result in a significant reduction in the production flow capacity of the created fracture. If an attempt is made to increase the stimulation fluid viscosity to a level whereby
fluid loss can be controlled without significant quantities of particulate fluid loss additives, the friction pressures resulting from the higher viscosity fluid limit the pumping rate and diminish the ability to produce a desired fracture length.

[0008] We have now devised a method of controlling fluid loss in a permeable formation, whereby the problems associated with the prior art are reduced or overcome.

[0009] According to the present invention, there is provided a method of controlling fluid loss in a permeable formation penetrated by a wellbore, which method comprises mixing an aqueous fluid containing at least a trace amount of at least one divalent cation, with a polymer derivative which polymer derivative comprises a reaction product made by the reaction of either (i) a hydroxyalkyl cellulose having a molar substitution of from 1.5 to 3, the alkyl being ethyl or propyl, or (ii) a graft copolymer of guar and hydroxypropyl guar, with (iii) a vinyl phosphonic acid in the presence of a redox system at a temperature of from 20 to 60°C, the polymer being present in an amount sufficient to viscosity said aqueous fluid, and a cross linking activator comprising a Lewis base or a Bronsted-Lowry base present in a sufficient amount to initiate crosslinking of at least a portion of said polymer derivative in said aqueous fluid to produce a crosslinked gel; shearing the crosslinked gel such that it breaks into particles having an average diameter in the range of from 1 mm to 10 mm; slurring the gel particles with an aqueous fluid having a density similar to the density of the gel particles whereby a suspension of the particles is produced; introducing the suspension into said wellbore and into contact with a face of said permeable formation; and producing a filter cake comprised of said particles of gel upon the face of said permeable formation whereby fluid loss to said formation through said filter cake is substantially reduced.

[0010] A particularly preferred fluid loss agent is prepared from certain graft copolymers of hydroxyethyl or hydroxypropyl cellulose, prepared by a redox reaction with vinyl phosphonic acid monomers or polymers and hydroxyethyl or hydroxypropyl cellulose which is crosslinked by the addition of a Lewis base of Bronsted-Lowry base or mixture of such bases to an aqueous solution of the graft copolymer. Certain graft copolymers of guar and hydroxypropyl guar prepared by a redox reaction with vinyl phosphonic acid monomers or polymers also may be crosslinked and utilized in a suitable manner. The crosslinked polymer gel then is sheared in any suitable manner to yield a particulate gel which is slurred or suspended in an aqueous fluid having a density similar to that of the gel.

[0011] To facilitate a complete understanding of the present invention, the following description will be directed to the use of particularly preferred polymers for use as the gelling agent. It is to be understood that such specificity is not to be construed as a limitation upon the scope of the present invention but is intended to aid in an understanding of the method.

[0012] A detailed description of the methods of manufacture of the preferred graft polymers used in the present invention is given in U.S. A-5,304,620, to which reference should be made for further details.

[0013] An aqueous liquid is used to solubilize the polymer or copolymer of the present invention. The term "aqueous liquid" is used hereafter to mean any liquid containing sufficient water to at least partially hydrate the copolymer and result in an increase in the viscosity of the fluid. Aqueous liquids used in oilfield operations normally contain sodium chloride, potassium chloride, calcium chloride, sodium bromide and other bromides, ammonium chloride, tetramethylammonium chloride or the like to weight the fluid or inhibit the swelling of clays generally found in subterranean formations. The pH of the aqueous liquid must be compatible with the selected crosslinking agent and must not adversely affect the hydration of the copolymer.

[0014] The term "vinyl phosphonic acid or VPA" as used herein includes not only the acid monomers and polymers but is also intended to include the aqueous soluble salts such as sodium or potassium vinyl phosphate and the like whereby a vinyl phosphonate ion is formed upon dissolution in an aqueous fluid as well as mono esters of vinyl phosphonic acid.

[0015] The cellulose derivative of this invention is preferably a hydroxyalkyl cellulose having a hydroxyalkyl molar substitution from about 1.5 to about 3.0. Molar substitution is defined as the average number of moles of a substituent group present per anhydroglucose unit of the cellulose material. The alkyl group is selected from the group of ethyl, propyl and mixtures thereof. The preferred hydroxyalkyl cellulose is hydroxyethyl cellulose (HEC) having a molar substitution in the range of about 1.8 to about 2.5. Preferably in this invention, the hydroxyalkylation of the cellulose is performed in a separate reaction. Hydroxyethyl cellulose is usually formed by reacting ethylene oxide with cellulose under extreme alkaline conditions and is available commercially.

[0016] The copolymers of the present invention are rendered crosslinkable by grafting monomers comprising a vinyl phosphonic acid to the cellulose derivative. The monomers have the reactive CH₂=C- moiety that is believed to enable the monomer to attach to the cellulose derivative.

[0017] Typically, graft copolymerizations are carried out in aqueous media wherein the polymer is dissolved or dispersed. The ratio of cellulose derivative to aqueous medium ranges from about 1 gram per 100 ml to about 1 gram per 2 ml. The preferred ratio is from about 1 gram per 6 ml to 1 gram per 4 ml. The ratio of cellulose derivative to grafting VPA monomer ranges from about 3 grams per 1 ml to about 25 grams per 1 ml. The preferred ratio is from about 6 grams per 1 ml to about 16 grams per 1 ml.

[0018] In one embodiment of the present invention, the crosslinkable copolymers are prepared by reacting certain
vinyl monomers having a crosslinkable substituent with a cellulose derivative using a redox system comprising, for example, the reaction product of hydrogen peroxide with a ferrous salt. The generalized redox reaction is believed to be represented by the formula:

\[ \text{H}_2\text{O}_2 + \text{Fe}^{2+} \rightarrow \text{HO}^* + \text{HO}^- + \text{Fe}^{3+} \]

and the generalized initiation reaction is believed to be represented by the general formula:

\[ \text{RCH}_2\text{OH} + \text{HO}^* \rightarrow \text{H}_2\text{O} + \text{RCH}_2\text{O}^* \]

[0019] An advantage of this initiator is that radical production occurs at a reasonable rate over a wide temperature range whereby reactions can be carried out at room temperature, if desired. The free radical produced on the cellulose derivative initiates polymerization with the vinyl group of the monomer to produce the graft copolymer.

[0020] Typically, the graft copolymerization is carried out in aqueous media wherein the polymer is partially dissolved or dispersed. Copolymers were prepared in acetone/water mixtures containing from about 55 to about 90% acetone. Reactions were carried out in a 1 liter kettle with a stirrer or a 1 liter jar at a temperature of from about 20 to about 60°C. The ratio of cellulose derivative to aqueous medium ranges from about 1 gram per 100 ml to about 1 gram per 2 ml. The preferred ratio is from about 1 gram per 2 to 5 ml. The ratio of cellulose derivative to grafting VPA monomer ranges from about 5 to about 40 grams per 1 gram of monomer. The preferred ratio is from about 6 to about 16. It is to be understood that the ranges set forth above are merely exemplary and that other temperatures, concentrations and the like maybe utilized to prepare the reaction product.

[0021] The polymerization reaction of this embodiment of the invention is chemically initiated by a redox system comprising the reaction product of hydrogen peroxide with a ferrous salt. Ferrous ions may be provided, for example, by salts such as ferrous ammonium sulfate, ferrous chloride, ferrous sulfate, ferrous acetate, ferrous oxalate, ferrous acetylacetonate and the like. A preferred source of ferrous ions is ferrous ammonium sulfate. Alternatively, other commonly used metal ion reductants may be utilized in place of the ferrous ions to generate the free radicals necessary to effect grafting and other forms of hydrogen peroxide such as t-butylhydroperoxide may be used.

[0022] The initiator may be added slowly to the reaction material over a time period of about 30 to 90 seconds or longer. Reaction times vary from about 15 minutes to about 4 hours depending upon the reaction conditions or the particular grafting monomer. Grafting reaction efficiency (% of monomer grafted) is generally less than about 75%. After the reaction is complete, the polymerization product is washed with acetone, filtered and dried.

[0023] In a preferred method of effecting the graft copolymerization, the graft polymer product is retained in a substantially storage stable slurry form. Typically, the media comprises a polyglycol, such as polypropylene glycol having molecular weights up to about 1000 such as "PPG-250" to "PPG-1000" polyglycol from Texaco Chemical Co., various polyethylene glycols and homopolymers of 1,2 butylene oxide having a molecular weight of from about 200 to about 400 which is present in an amount of from about 70 to about 95 percent by weight of the media and the remainder generally comprises water. The media also may comprise tetramethylammonium chloride in a similar amount or in admixture with a polyglycol. In a preferred embodiment the polyglycol comprises from about 85 to 92 percent by weight of the media. Reactions were conducted in a 5 liter kettle with a stirrer at a temperature of from about 20 to about 60°C. The ratio of cellulose derivative to media ranges from about 1 gram per 100 ml to about 1 gram per 2 ml. The preferred ratio is from about 1 gram per 2 to 5 ml. The reaction media also may include a quantity of a dispersant or thixotrope such as alkylation quaternary ammonium montmorillonite ("CLAYTONE AF" thixotrope from E.C.C. America, Inc.) or dimethylidioctoammonium chloride to facilitate dispersion of the polymer in the media and improve suspension properties. The grafting reaction is performed as previously described using an appropriate redox system such as, for example, the ferrous salt with a source of peroxide. Since the metal ions are not removed from the product by washing as when a dry product is formed, a sequestrant for the metal ions may be added to the slurry at the conclusion of the reaction. The polymerization product has been found to remain readily dispersible or suspended in the slurry form over a period of time to facilitate storage and handling.

[0024] Graft copolymers of the present invention solubilize in aqueous liquids and substantially increase the viscosity of aqueous liquids. The viscosity of the copolymer solution is further increased with the addition of a selected crosslinking activator or agent which effects an initiation of a crosslink interaction. Preferred crosslinking activators or agents in accordance with this invention comprise Bronsted-Lowry or Lewis bases. Suitable compounds are believed to include, for example, calcium oxide, magnesium oxide and compounds selected from mon, di and trialkanol amines such as triethanolamine, sodium hydroxide, potassium hydroxide, ammonia, various cocoaimes such as Bis(2-hydroxyethyl) cocoaime, various pentamines such as tetraethylenepentamine, and various other water soluble amines, such as propyl/diethanolamine, triethyamine, various water soluble borates such as the commercially available product
"POLYBORB" an admixture of boric acid and borate salts, from U.S. Borax and the like in the presence of a divalent cation such as calcium or magnesium which is present in at least a trace amount and which may be present in the aqueous liquid utilized to hydrate the copolymer or added as an additional component to the aqueous liquid. Such compounds generally are substantially free of polyvalent metal ions, that is, metal ions having more than one valence state. A particularly preferred crosslinking agent is magnesium oxide.

While the specific mechanism by which the crosslinking occurs is unknown, it is believed that the crosslink is formed through the phosphorus moiety in the graft polymer which is activated by the presence of the Lewis or Bronsted-Lowry base.

While the following description will be directed to the use of magnesium oxide as a crosslinking activator or agent, it is to be understood that the described method would apply generally to any of the other suitable crosslinking agents of the present invention. A base gel is prepared by hydrating the previously described graft copolymer of hydroxyethyl cellulose or hydroxypropyl cellulose, in an aqueous fluid at a pH in the range of from about 0.1 to about 3.5. The graft copolymer can be admixed with the aqueous fluid in an amount of from about 10 to about 175 pounds per 1000 gallons (1.2 to 20.9 kg m⁻³) of fluid. After the gelling agent has been substantially hydrated, the base gel is admixed with a quantity of magnesium oxide in an amount of from about 10 to about 40 pounds per 1000 gallons (1.2 to 4.8 kg m⁻³) of base gel. In a preferred method of addition, the magnesium oxide is added as an aqueous slurry to facilitate handling of the material.

The crosslinking of the graft copolymer solution may be accomplished in any suitable vessel or apparatus such as, for example, a vertical paddle blender or horizontal ribbon blender or the like. The crosslinked gelled fluid then is suitably sheared to effect formation of discrete particles of the gelled fluid. In one embodiment, the fluid is sheared by passage through a perforated plate at an elevated pressure whereupon the gel breaks into generally small cylindrical shaped particles. Alternatively, the gelled fluid may be passed through a crossed network of thin wires within a circular or square tube to form various geometries of gel particles. Generally, the particles of crosslinked gelled fluid will have an average diameter in the range of from about 1 mm to about 10 mm if of a generally spherical nature and an average diameter in the range of from about 1 mm to about 10 mm and a length of from about 1 mm to about 50 mm if of a cylindrical nature. The particles of crosslinked gelled fluid then are slurried or suspended in an aqueous fluid having a density similar to that of the crosslinked gelled fluid. The aqueous suspension fluid may comprise substantially any of the previously described aqueous fluids and may have a density either greater than or less than that of the crosslinked gelled fluid. Preferable the aqueous suspension fluid has a density within ± 0.5 of the density of the crosslinked gelled fluid to facilitate formation of the suspension and transport of the particles.

The particulate slurry then may be utilized to provide fluid loss control in a wellbore penetrating a subterranean formation by introduction into the wellbore. A filter cake of the suspended particles of gel is formed upon the face of the formation as fluid loss occurs to the formation. The fluid loss control can be effected when used with drilling, completion and stimulation fluids. The quantity of particulate gel used in accordance with the different aspects of the present invention is that amount which is sufficient to achieve a desired level of fluid loss control. The quantity will depend, to some extent, upon the permeability of the formation and formation temperature, or well as the desired level of fluid loss control.

A beneficial aspect of the use of the graft copolymers described herein is the ability of the particulate gel to be readily broken to a low viscosity fluid to facilitate cleanup of the formation face contacted by the particulate and minimize any potential permeability-loss problem resulting from use of the fluid loss control agent. The crosslinked gel particulate may be readily broken by contacting the particulate with an acidic fluid. The acidic fluid can comprise substantially any fluid having a pH less than that of the crosslinked gel whereby upon contact with the gel the pH of the gel is reduced. Examples of such fluids and compounds include, but are not limited to, hydrochloric acid solutions, organic acids such as fumaric acid, acetic acid, citric acid, polyhydroxyacetic acid, encapsulated forms of acid generating compounds or acids and the like.

The particulate slurry also can contain substantially any other conventional additives, such as for example, gel stabilizers, breakers, clay stabilizers, bactericides, fluid loss additives, surfactants, weighting agents such as hematite, barite or calcium carbonate and the like. The selection of such additives is well within the skill of those experienced in the art.

The following examples are provided to illustrate the utility of the method of the present invention.

**EXAMPLE I**

The following tests were performed to compare the effectiveness of a high viscosity gel to a particulate gel slurry in controlling fluid loss through a permeable formation. Berea sandstone disks of approximate 100 mD permeability, 4.48 cm in diameter and 0.72 cm thick were placed into high pressure fluid loss cells and the temperature of the cell was raised to 120°F (48.9°C). A solution of 2% NH₄Cl brine, in an amount of about 300 ml, was flowed through each disk to saturate the sample by application of 100 psi (14.5 x 10⁻⁴ Pa) nitrogen gas pressure. A gel solution was
prepared by admixing the equivalent of 120 lb of polymer per 1000 gallons (14.4 kg m\(^{-3}\)) of brine of the vinyl phosphonic acid grafted hydroxyethyl cellulose into a 9.0 lb per gallon density CaCl\(_2\) completion brine. The equivalent of 19 \(\ell\) per 3800 \(\ell\) (5 gal per 1000 gallons) of brine of 37% HCl also was added to the completion brine to facilitate hydration of the copolymer. After 30 minutes, a slurry of MgO in completion brine equivalent to a concentration of 30 lb per 1000 gallons (3.6 kg m\(^{-3}\)) of brine was added to the hydrated copolymer solution. After crosslinking, the mixture was placed on one Berea core in a heated fluid loss cell. The cell was closed and 100 psi (14.5 x 10\(^{-3}\) Pa) nitrogen gas pressure was applied to the sample in the cell. The fluid loss versus time is set forth in Table I. A second gel sample was prepared as indicated above with the additional steps of passing the gel through a perforated plate having 1/8 inch (0.3175 cm) diameter holes using 100 psi (14.5 x 10\(^{-3}\) Pa) nitrogen gas pressure and mixing the extruded gel particles with a quantity of the 9.0 lb per gallon (1078 kg m\(^{-3}\)) density CaCl\(_2\) brine at a ratio of 4 parts by volume gel to 1 part by volume brine. This slurry was then placed in the fluid loss cell on a Berea core and the fluid loss test was repeated. The results also are set forth in Table I.

### TABLE I

<table>
<thead>
<tr>
<th>Time, Minutes</th>
<th>Fluid Loss Versus Time Volume, ml</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Polymer Sample</td>
</tr>
<tr>
<td>1</td>
<td>1.0</td>
</tr>
<tr>
<td>2</td>
<td>1.5</td>
</tr>
<tr>
<td>3</td>
<td>2.0</td>
</tr>
<tr>
<td>4</td>
<td>2.0</td>
</tr>
<tr>
<td>5</td>
<td>2.0</td>
</tr>
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<td>6</td>
<td>2.5</td>
</tr>
<tr>
<td>7</td>
<td>2.5</td>
</tr>
<tr>
<td>8</td>
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</tr>
<tr>
<td>9</td>
<td>2.5</td>
</tr>
<tr>
<td>10</td>
<td>2.8</td>
</tr>
<tr>
<td>15</td>
<td>3.0</td>
</tr>
<tr>
<td>30</td>
<td>4.0</td>
</tr>
</tbody>
</table>

[0033] The above data clearly illustrates the ability of the particulate gel to provide fluid loss control.

### EXAMPLE II

[0034] To illustrate the ease with which the particulate gel of the present invention can be pumped through tubing and into a formation, the following tests were performed. A gel sample was prepared as in Example I. The viscosity of the gel was measured using a Brookfield “LVT” viscometer equipped with a helical path motion device and a size F “T” spindle at 3 RPM. The viscosity measurement was made at room temperature. The viscosity of a particulate gel sample prepared as in Example I using dilution ratios of polymer to brine of 1:1, 2:1, 3:1 and 4:1 also was determined. The results of the viscosity measurements are set forth below in Table II.

### TABLE II

<table>
<thead>
<tr>
<th>SAMPLE</th>
<th>VISCOSITY, cps x 10(^6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100% Crosslinked Gel</td>
<td>Not measurable</td>
</tr>
<tr>
<td>1:1 polymer/brine ratio</td>
<td>1.20</td>
</tr>
<tr>
<td>2:1 polymer/brine ratio</td>
<td>4.20</td>
</tr>
<tr>
<td>3:1 polymer/brine ratio</td>
<td>7.56</td>
</tr>
<tr>
<td>4:1 polymer/brine ratio</td>
<td>9.99</td>
</tr>
</tbody>
</table>

[0035] Since friction pressure in a pipe is proportional to the viscosity of a fluid pumped through the pipe, the data
clearly illustrates that friction pressure and hence pumping pressures can be controlled through appropriate dilution of the particulate gel slurry.

5 Claims

1. A method of controlling fluid loss in a permeable formation penetrated by a wellbore, which method comprises mixing an aqueous fluid containing at least a trace amount of at least one divalent cation, with a polymer derivative which polymer derivative comprises a reaction product made by the reaction of either (i) a hydroxyalkyl cellulose having a molar substitution of from 1.5 to 3, the alkyl being ethyl or propyl, or (ii) a graft copolymer of guar and hydroxypropyl guar, with (iii) a vinyl phosphonic acid in the presence of a redox system at a temperature of from 20 to 60°C, the polymer being present in an amount sufficient to viscosity said aqueous fluid, and a cross linking activator comprising a Lewis base or a Bronsted-Lowry base present in a sufficient amount to initiate crosslinking of at least a portion of said polymer derivative in said aqueous fluid to produce a crosslinked gel; shearing the crosslinked gel such that it breaks into particles having an average diameter in the range of from 1 mm to 10 mm; slurring the gel particles with an aqueous fluid having a density similar to the density of the gel particles whereby a suspension of the particles is produced; introducing the suspension into said wellbore and into contact with a face of said permeable formation; and producing a filter cake comprised of said particles of gel upon the face of said permeable formation whereby fluid loss to said formation through said filter cake is substantially reduced.

2. A method according to claim 1, wherein the crosslinked gel is sheared by passage through a perforated plate at a sufficient rate and pressure to cause formation of said particles.

3. A method according to claim 1 or 2, wherein the crosslinking activator comprises at least one of magnesium oxide; a mono, di or trialkanamide; calcium oxide; sodium hydroxide; potassium hydroxide; ammonia; a cocooamine; a pentamine; an alkylidethanolamine; a mixture of boric acid and borate salts and diethyamine.

4. A method according to claim 1, 2 or 3, wherein the reaction to produce the polymer derivative is effected in a reaction medium comprising at least one of tetramethyl ammonium chloride, polyethylene glycol and polypropylene glycol.

5. A method according to claim 1, 2, 3 or 4, wherein the redox system comprises a peroxide and a metal ion reductant.

6. A method according to claim 5, wherein the metal ion reductant comprises a source of ferrous ions.

7. A method according to any of claims 1 to 6, wherein said gel particles have an average length of from 1 to 50 mm.

Patentansprüche

1. Methode zur Bekämpfung des Fluidverlustes in einer durchlässigen, durch ein Bohrloch durchdrungenen Formation, welche Methode folgendes umfaßt: Mischen eines mindestens eine Spurenmengen mindestens eines zweiwertigen Kations enthaltenden wässrigen Fluids mit einem Polymerderivat, das ein Reaktionsprodukt umfaßt, das durch Reaktion entweder (i) einer Hydroxyalkylcellulose mit einer molaren Substitution von 1.5 bis 3 - wobei es sich bei dem Alkyl zum Ethyl oder Propyl handelt - oder (ii) eines Ppropfopolymer von Guar und Hydroxypropylguar mit (iii) einer Vinylphosphonsäure in Gegenwart eines Redoxsystems bei einer Temperatur von 20 bis 60°C hergestellt wird, wobei das Polymer in einer Menge vorliegt, die ausreicht, um die Viskosität des wässrigen Fluids zu erhöhen, und einem Vernetzungsaktivator, der eine Lewis-Base oder eine Bronsted-Lowry-Base umfaßt, die in einer Menge vorliegt, die ausreicht, um die Vernetzung mindestens eines Teils des Polymereinheits in dem wässrigen Fluid unter Bildung eines vernetzten Gels einzuleiten; Unterziehen des vernetzten Gels einer Scherbeanspruchung derart, daß es in Teilchen mit einem Durchmesserdurchmesser im Bereich von 1 mm bis 10 mm zerfällt. Aufschlamm der Gelteilchen mit einem wässrigen Fluid, die eine Dichte ähnlich der der Gelteilchen aufweist, wobei eine Suspension der Teilchen gebildet wird; Einführen der Suspension in das Bohrloch und in Berührung bringen mit einer Stirnseite der durchlässigen Formation, und Herstellen eines Filterkuchens, der aus den Gelteilchen auf der Stirnseite der durchlässigen Formation besteht, wobei der Verlust an Fluid an die Formation durch den Filterkuchen wesentlich reduziert wird.

2. Methode nach Anspruch 1, bei der das vernetzte Gel einer Scherbeanspruchung ausgesetzt wird beim Durchgang
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durch eine durchlöcherte Platte mit einer Geschwindigkeit und einem Druck, die zur Bildung der Teilchen ausreich.

3. Methode nach Anspruch 1 oder 2, bei der der Vernetzungsaktivator mindestens eine der folgenden Substanzen umfaßt: Magnesiumoxid; ein Mono-, Di- oder Trialkanolamin; Calciumoxid; Natriumhydroxid; Kaliumhydroxid; Ammoniak; ein Kokosamin; ein Pentamin; ein Alkylidithanolamin; eine Mischung von Borsäure und Boratsalzen und Diethylamin.

4. Methode nach Anspruch 1, 2 oder 3, bei der die Reaktion zur Bildung des Polymerderivats in einem Reaktionsmedium durchgeführt wird, das mindestens eine der folgenden Substanzen umfaßt: Tetramethylammoniumchlorid, Polyethylen glykol und Polypropylen glykol.

5. Methode nach Anspruch 1, 2, 3 oder 4, bei der das Redoxsystem ein Peroxid und ein Metallionenreduziermittel umfaßt.


7. Methode nach einem der Ansprüche 1 bis 6, bei der die Teilchen eine durchschnittliche Länge von 1 bis 50 mm aufweisen.

Revendications

1. Un procédé de régulation des pertes de fluide dans une formation perméable traversée par un trou de sondage, dit procédé comportant le mélange d’un fluide aqueux contenant au moins une trace d’eau moins un cation divalent, avec un dérivé polymérique qui comporte un produit de réaction élaboré par la réaction de soi (i) une hydroxyl cellulose ayant une substitution molaire de 1,5 à 3, l’étyle étant étyle ou propyle, soit (ii) un copolymère greffé de guar et d’hydroxypropyl guar, avec (iii) un acide phosphonique vinylique en présence d’un système rédox à une température de 20 à 60°C, le polymère étant présent en une quantité suffisante pour viscosifier ledit fluide aqueux, et un activateur de rétication qui comprend une base de Lewis ou une base de Bronsted-Lowry présente en une quantité suffisante pour amorcer la rétication d’eau moins une partie dudit dérivé polymérique dans le fluide aqueux pour produire un gel réticulé; cisaillé du gel réticulé de façon telle qu’il se désintègre en particules ayant un diamètre moyen dans la plage de 1 mm à 10 mm; mise en boue aqueuse des particules de gel afin de produire une suspension de particules, introduction de ladite suspension dans le trou de sondage et en contact avec une face de ladite formation perméable, et production d’un gâteau de filtration composé desdites particules de gel sur la face de ladite formation perméable, ce qui réduit sensiblement les pertes de fluide vers ladite formation par le biais dudit gâteau de filtration.

2. Un procédé selon la revendication 2, selon lequel le gel réticulé est cisaillé par passage à travers une plaque perforée à une vitesses et à une pression suffisantes pour causer la formation desdites particules.

3. Un procédé selon la revendication 1 ou 2, selon lequel l’activateur de réticulation comporte au moins l’une des molécules suivantes : oxyde de magnésium ; une mono, di ou trialcanolamine ; un oxyde de calcium ; un hydroxyde de sodium (soude caustique) ; un hydroxyde de potassium (potassium caustique) ; ammoniac ; une cocoamine ; une alkylidethanolamine ; un mélange d’acide borique et de sels de borate et diéthylamine.

4. Un procédé selon la revendication 1, 2 ou 3, selon lequel la réaction de production du dérivé polymérique est menée dans un milieu de réaction comportant au moins l’une des molécules : chlorure d’ammonium tétraméthylène, polyéthyléneglycol et polypropylène glycol.

5. Un procédé selon la revendication 1, 2, 3 ou 4, selon lequel le système rédox comporte un peroxyde et un ion de métal réducteur.

6. Un procédé selon la revendication 5, selon lequel l’ion de métal réducteur comporte une source d’ions ferreux.

7. Un procédé selon l’une quelconque des revendications 1 à 6, selon lequel lesdites particules de gel ont une longueur moyenne de 1 à 50 mm.