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(54) Title: METHOD FOR TREATING OIL-CONTAINING AQUEOUS MIXTURES WITH CATION EXCHANGE RESIN

(57) Abstract: A method for treating an oil-containing aqueous mixture comprising the step of passing the mixture (1) through a media (6) comprising a bead-form cation exchange resin combined with a cationic surfactant, wherein the resin includes a sulfonated crosslinked copolymer matrix having a rough surface characterized by having a frequency of at least 5 peaks and valleys per sample surface area (283 μm x 212 μm), where the difference between the average height of the 5 highest peaks and the 5 lowest valleys is at least 6 μm.

[Continued on next page]
METHOD FOR TREATING OIL-CONTAINING AQUEOUS MIXTURES
WITH CATION EXCHANGE RESIN

FIELD

The invention relates to methods for separating oil from aqueous mixtures using ion
exchange media.

INTRODUCTION

Emulsified oily aqueous mixtures are produced in a wide variety of industrial processes
including those used in the food, petroleum, petrochemical and metallurgical industries. Emulsified
oil wastewater is hazardous for aquatic environments and generally requires treatment prior to
disposal or re-use. One treatment approach described by Y.B. Zhou involves the use of coalescence;
see: 1) Emulsified oil wastewater treatment using a hybrid modified resin and activated carbon
system, Separation and Purification Technology (2008), 63, 400-406; 2) Study on resin modified by
quaternary ammonium salt for emulsified oil wastewater treatment, China Water Wastewater
(2008), 24, 60-63; 3) Modified resin coalesce for oil-in-water emulsion treatment: effect of
resin, Separation and Purification Technology (2010), 75, 266-272. See also: B.W. Yang et al., 1)
continues to search for improved methods and media for removing oil from aqueous mixtures
including industrial waste water.

SUMMARY

In one embodiment the invention includes a method for treating an oil-containing aqueous
mixture by passing the mixture through a media including a bead-form cation exchange resin
combined with a cationic surfactant. The resin includes a sulfonated crosslinked copolymer matrix
with a rough surface characterized by having at least 5 peaks and valleys per sample surface area (283
um x 212 um) where the difference between the average height of the 5 highest peaks and the 5 lowest
valleys is at least 6 um.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A is a schematic representation of a coalescer as used in the Examples.

Figures 2a-d are micrographs of bead-form cation exchange resins as described in the
Example section.
DETAILED DESCRIPTION

The invention includes a method for separating oil from an aqueous mixture. Neither the type (e.g., mineral, organic) nor source (e.g., animal, vegetable, petrochemical) of oil is particularly limited. The invention finds particular utility in treating oil-containing aqueous mixtures generated or used in the food, petroleum, petrochemical and metallurgical industries.

In a preferred embodiment, the oil-containing aqueous mixture is passed through a media bed or column that serves as a coalescer. A schematic view of representative coalescer is generally shown at 10 in Figure 1 including a tank equipped with an oily feed water intake (1), a treated water outlet (2), a recovered oil outlet (3), a wash water inlet (4), general outlet (5) and packed media section (6). In operation, an oil-containing aqueous feed mixture flows into the coalescer by way of intake (1) and passes through the media (6). As the feed mixture passes upward through the media (6), oil is adsorbed onto the media surface and progressively grows to form droplets (7). The oil droplets (7) float to the top of the coalescer to form an oil layer (8) which may be removed, e.g., by way of skimming. Treated liquid may be recovered at a point below the oil layer, e.g., by outlet (3).

Alternative configurations and fluid flow patterns may also be used. The recovered water may be further treated, e.g., by way of filtration, passing through a bed of adsorbent (e.g., OPTIPORE L493 available from The Dow Chemical Co.), passing through another bed of ion exchange resin, etc.

The media includes a cation exchange resin provided in bead form. The resin preferably includes crosslinked copolymer matrix derived from polymerizing a monomer mixture including a monovinyldene monomer such as styrene and a crosslinker such as divinylbenzene, e.g., by suspension polymerization of a finely divided organic phase comprising styrene along with other optionally monovinyldene monomers, crosslinking monomers including divinylbenzene, a free-radical initiator and optionally a phase-separating diluent. While the crosslinked copolymer may be macroporous or gel-type, gel-type copolymers are preferred. The terms “gel-type” and “macroporous” are well-known in the art and generally describe the nature of the copolymer bead porosity. The term “macroporous” as commonly used in the art means that the copolymer has both macropores and mesopores. The terms “microporous,” “cellular,” “gel” and “gel-type” are synonyms that describe copolymer beads having pore sizes less than about 20 Angstroms Å, while macroporous copolymer beads have both mesopores of from about 20 Å to about 500 Å and macropores of greater than about 500 Å. Gel-type and macroporous copolymer beads, as well as their preparation are further described in US 4,256,840 and US 5,244,926 – the entire contents of which are incorporated herein by reference. The crosslinked copolymer resin beads preferably have a median bead diameter from 100 to 2000 microns. The beads may have a Gaussian particle size distribution but preferably have a relatively uniform particle size distribution, i.e., “monodisperse” that is, at least 90 volume percent of the beads have a particle diameter from about 0.9 to about 1.1 times the volume average
particle diameter.

Once formed, the crosslinked copolymer resin beads are sulfonated with a sulfonating agent, e.g. sulfuric acid, chlorosulfonic acid or sulfur trioxide. The sulfonation reaction is preferably conducted at elevated temperature, e.g. 100 to 150°C without a solvent. An applicable sulfonation technique is described in US 2005/0014853. See also: US 6228896, US 6750259, US 6784213, US 2002/002267 and US 2004/0006145. Sulfonation of the resin in the absence of a solvent is believed to impart a rough or dimpled surface to the resin bead. By way of illustration, Figures 1a-d are micrographs of two comparable ion exchange resins. The resin shown in Figs. a) and b) was sulfonated using a traditional solvent (EDC) whereas the resin shown in Figs c) and d) was sulfonated in the absence of a solvent.

In a preferred embodiment, the surface of the resin is characterized by including a frequency of at least 5 (preferably at least 10) peaks and valleys per sample surface area (283 um x 212 um). The difference between the average height of the 5 highest peaks and the 5 lowest valleys \( (S_{50\%}) \) value is preferably at least 6 μm, more preferably at least 8 μm and even more preferably at least 10 μm. In another embodiment, the resin has a surface area ratio \( (S_{5\%}) \) of from 0.9% to 2.4%, wherein the “surface area ratio” refers to the increment of the interfacial surface area relative to an area of the projected (flat) x,y plane, (i.e. for a flat surface, the surface area of the x-y plane are the same and \( S_{5\%} = 0\% \)). Surface characterization may be performed using a confocal laser scanning microscope (CLSM), preferably one using a scanning violet (408 nm) laser light source for high resolution confocal surface profiling. A preferred resin is AMBERLITE SR1L available from The Dow Chemical Company.

Once sulfonated, the resulting cation exchange resin is combined with a cationic surfactant. In a preferred embodiment the surfactant is loaded onto the resin by preparing an aqueous solution of the surfactant and passing the solution through a bed of the cation exchange resin. The resin may also be soaked in such a solution. The resin is preferably then rinsed with water prior to use. The surfactant is preferably loaded on the resin at ratio of 5% to 99% of the total exchange capacity of the resin, (i.e. the positively charged moiety of the surfactant associates with the negatively charged sulfonate functional groups of the cation exchange resin). The cationic surfactant includes a functional group selected from at least one of the following: i) an amine/ (primary, secondary or tertiary) and ii) a quaternary ammonium, iii) pyridinium, iv) quinoline, v) bis-quaternary ammonium, vi) sulfonium, vii) phosphonium and viii) arsonium. Applicable surfactants include those represented by the following formulae:
wherein X is selected from a halogen and R₁ through R₁₀ are independently selected from hydrogen and hydrocarbys having from 1 to 20 carbon atoms with the proviso that at least one of R₁ through R₁₀ are independently hydrophobic moiety of alkyl, alkylarylalkyl, alkoxyalkyl, alkylaminoalkyl and alkylamidoalkyl, where alkyl represents the carbon chain length from a to b and which may branched or straight chain and which may be saturated or unsaturated. Preferred species including: dodecyl dimethyl benzyl ammonium bromide, hexadecyltrimethylammonium bromide, hexadecylpyridinium bromide, tetrabutylammonium bromide and cetyltrimethyl-ammonium bromide.

EXAMPLES

To better illustrate the invention, two test runs were completed using the same oily feed mixture, coalescer and operating conditions but using two different cation exchange resins. In both test runs, the resins were combined with the same cationic surfactant according to the same methodology. The experimental set up is schematically illustrated in Figure 1.

Coalescer: 1600 ml resin packing volume
Feed mixture: 140 mg/L of oil at pH of approx. 3 and a temp. of approx. 36°C
Operating conditions: Volumetric flow rate of 8BV/h
Cationic surfactant: Dodecyl dimethyl benzyl ammonium bromide
Resin of Test Run 1: AMBERLITE 120 Na
Resin of Test Run 1: AMBERLITE SR1L
The first test run was conducted using AMBERLITE 120 Na brand cation exchange resin available from The Dow Chemical Company. This resin has a crosslinked, styrene-divinylbenzene copolymer matrix that has been sulfonated with H₂SO₄ using a solvent (ethylene dichloride). The second test run was conducted using AMBERLITE SR1L brand cation exchange resin also available from The Dow Chemical Company. This resin is substantially similar to AMBERLITE 120 Na; however, this resin was sulfonated using H₂SO₄ without solvent. The surfaces of both cation exchange resins were analyzed (prior to loading with cationic surfactant) with a confocal laser scanning microscope (CLSM) using a scanning violet laser (408 nm) light source for high resolution confocal surface profiling. Combined with an additional white light source, the system provided simultaneous color, laser intensity, and height information to generate high-resolution images. The CLSM was operated with a 1 nm z-resolution and 130 nm spatial resolution providing SEM-like images with a large 7mm through focus range. Figure 1 a) and b) are micrographs of AMBERLITE 120 Na and Figure 1 c) and d) are micrographs of AMBERLITE SR1L.

During each test run, treated water samples were collected at regular intervals and residual oil concentrations were determined using UV spectroscopy. The oil removal rate of the second test run increased over 20% from the first test run. Such higher oil removal rates permit greater flexibility in coalescer design and use of higher flow rates along with provided more options for further downstream treatment.
CLAIMS

1. A method for treating an oil-containing aqueous mixture comprising the step of passing the mixture through a media comprising a bead-form cation exchange resin combined with a cationic surfactant, wherein the resin comprises a sulfonated crosslinked copolymer matrix having a rough surface characterized by comprising a frequency of at least 5 peaks and valleys per sample surface area (283 µm x 212 µm), where the difference between the average height of the 5 highest peaks and the 5 lowest valleys is at least 6 µm.

2. The method of claim 1 wherein the surface of the resin is characterized by comprising a frequency of at least 10 peaks and valleys per sample surface area (283 µm x 212 µm).

3. The method of claim 1 wherein the cation exchange resin has a surface area ratio of from 0.9% to 2.4%.

4. The method of claim 1 wherein the resin is a gel-type.

5. The method of claim 1 wherein the crosslinked copolymer matrix is derived from polymerizing a monomer mixture comprising styrene or divinylbenzene.

6. The method of claim 1 wherein the cationic surfactant comprises a functional group selected from at least one of the following: i) an amine and ii) a quaternary ammonium iii) pyridinium, iv) quinoline, v) bis-quaterenary ammonium, vi) sulfonium, vii) phosphonium and viii) arsonium.

7. The method of claim 1 wherein the cationic surfactant is represented by the formula:

\[
\begin{align*}
R_1 & \quad R_2 \\
\text{N} & \quad X^- \\
R_3 & \quad R_4
\end{align*}
\]

wherein X is selected from a halogen and R₁, R₂, R₃ and R₄ are independently selected from hydrogen and hydrocarbyls having from 1 to 20 carbon atoms with the proviso that at least one of R₁, R₂, R₃ and R₄ is a hydrocarbyl.

8. The method of claim 1 wherein the resin comprises a sulfonated crosslinked copolymer matrix having a rough surface characterized by comprising a frequency of at least 5 peaks and valleys per sample surface area (283 µm x 212 µm), where the difference between the average height of the 5 highest peaks and the 5 lowest valleys is at least 8 µm.

9. The method of claim 1 wherein the resin comprises a sulfonated crosslinked copolymer matrix having a rough surface characterized by comprising a frequency of at least 5 peaks and valleys per sample surface area (283 µm x 212 µm), where the difference between the average height of the 5 highest peaks and the 5 lowest valleys is at least 10 µm.
IER-100na: Mag 100  IER-100na: Mag 500
IER-SR1L: Mag 100   IER-SR1L: Mag 500

Fig. 2a-d
**INTERNATIONAL SEARCH REPORT**

**A. CLASSIFICATION OF SUBJECT MATTER**

INV. B01D17/04 B01J39/20 C02F1/42

**ADD.**

According to International Patent Classification (IPC) or to both national classification and IPC.

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

B01D B01J C02F

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched.

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

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<td>YANBO ZHOU ET AL: &quot;Effect of quaternary ammonium surfactant modification on oil removal capability of polystyrene resin&quot;, SEPARATION AND PURIFICATION TECHNOLOGY, vol. 75, no. 3, 1 November 2010 (2010-11-01), pages 266-272, XP051422222, ISSN: 1383-5866, DOI: 10.1016/j.seppur.2010.08.015 paragraph 2.1 abstract</td>
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Further documents are listed in the continuation of Box C. See patent family annex.

* Special categories of cited documents:
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  * "E" - earlier application or patent but published on or after the international filing date
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**Date of the actual completion of the international search**

30 September 2014

**Date of mailing of the international search report**

03/11/2014

**Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-3040, Fax: (+31-70) 340-3016**

Authorized officer

Thibault, Valerie

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