A novel and cleaner bio-polymer Gum Karaya-based Silica nanocomposite fracturing fluid for high-temperature application

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Abstract
A novel polymer nanocomposite has been synthesized in the present study using a bio-polymer Karaya and inorganic SiO$_2$ nanoparticles via sonication for application as a fracturing fluid in stimulating oil and gas wells. Gum Karaya (KG), an acidic polysaccharide, has been investigated for its structural, morphological and rheological characteristics, and the results have been compared with prepared KG–SiO$_2$ nanocomposite at different SiO$_2$ dosing for improved structure, viscoelasticity, viscosity and temperature stability. Steady shear rheology test depicted pseudoplastic nature, while oscillatory study confirms gel characteristics and weak dependence of storage and loss moduli on frequency for all the prepared samples. Dominance of elasticity over viscous nature of the gel assures efficient solid transporting/suspending capability. Experimental results showed that the rheological characteristics of the synthesized gel is comparable with Guar and CMHPG over the shear and temperature ranges under study and suitable for temperatures above 150 °C and pressure conditions. Also, polymer breaking and sandpack regained permeability tests proved synthesized gels to be a cleaner with less residue as compared to the other two.

Keywords
Nanocomposite · Bio-polymer-Karaya · Thermal stability · Rheology · Gel breaking · Hydraulic fracturing

Introduction
The recent technological advancement in hydraulic fracturing (HF) has significantly enhanced the oil and gas production worldwide, especially in the USA. HF—a stimulation technique improves the reservoir permeability by utilizing fracturing fluid to create a fracture in the formation and to carry and place the proppant into the fracture. Conventionally, the water-soluble polymers like Guar and its derivatives like Carboxyl Methyl Hydroxyl Propyl Guar (CMHPG) in linear or metal ions-cross-linked form are being used as thickening agents to develop sufficiently high viscosity to fracture the formation under high-pressure and to carry the proppant to the fracture. It is desirable that these highly viscous gels should maintain their stability till the fracture opening and should be broken into a low viscosity fluid as soon as the proppant is placed into the fractures (Song et al. 2016; Yegin et al. 2016). However, partially or unbroken polymeric fluids generate residues upon breaking, causing a significant reduction in the effective permeability by clogging the pores of the formation and the proppant pack. The damage caused by these polymer fluids becomes more severe at high-temperature and high-pressure (HTHP) conditions where high polymer concentration is used to maintain the thermal stability. Application of nanotechnology for developing “smart nanofluids” could be adopted to solve this formation damage problem. These nanofluids have exceptionally high surface-to-volume ratio that provides the advantages of high thermal stability, elasticity, viscosity and wettability alteration to operate in HTHP conditions of the reservoir without increasing the polymer concentration (Cocuzza et al. 2011; Matteo et al. 2012; Song et al. 2016; Yegin et al. 2016).

In the last two decades, industrial sectors like pharmaceuticals, aerospace, automobile and oil and gas industry
have paid strong emphasis for the evolution of polymeric nanocomposite, bearing at least one reinforcing component with nanometer range size (Henrique et al. 2009). In the oil and gas industry, nanotechnology is being used for non-corrosive material development, downhole separation processes and for enhanced oil and gas productivity (Matteo et al. 2012). Polymer-nanocomposites are novel materials and have attracted many researchers in the recent years due to their hybrid and synergistic properties with more interfacial adhesion, which provides exceptional optical, electrical, thermal and mechanical properties (elastic modulus, strength and dimension stability) (Paul and Robeson 2008; Henrique et al. 2009). However, these materials are rarely used in the oil industries and very little is known about the shear influence on the combined soft polymer with rigid inorganic nanoparticles, which is important in evaluating their efficiency at the downhole conditions (Paul and Robeson 2008; Henrique et al. 2009). Motivated with the novel characteristics of nanopolymer composites and the problem faced in hydraulic fracturing with the polymers, a nanocomposite has been developed as a hydraulic fracturing fluid (HFF) with silicon dioxide nanoparticles and Gum Karaya polymer, and various properties related to HFF were studied and compared with conventionally used polymers in oil industry. A natural bio-polymer Gum Karaya (KG), also called gum Sterculia, has been used in the present study as polymer for preparing the nanocomposite, which has also been used in our previous study to prepare zirconium-cross-linked HFF and was found effective up to a temperature of 150 °C (Chauhan et al. 2019). KG is obtained from the large and mature Sterculia urens grown extensively throughout the Indian sub-continent in arid forest regions. The gum has found its potential as an emulsifier, thickener and stabilizer in the food and pharmaceutical industries (Abo-Shosha et al. 2008; Augusto et al. 2012; De Brito et al. 2005). Silicon dioxide offers promising support for the synthesis of bio-composite by developing active paramagnetic acid and base centers for mechanochemical reactions (Vorsina et al. 2011). In the present study, morphological, structural, sandpack regained permeability, steady state and dynamic rheology of aqueous solution of KG and KG–SiO₂ nanocomposite for improved properties have been studied and the results have been compared with conventional polymers like guar and CMHPG.

Material and methods

Materials

KG having molecular weight (MW) 950,000 Da and silicon dioxide (SiO₂) nanopowder having 99.5% purity and 15 nm average particle size were procured from Sigma-Aldrich, UK. CMHPG (MW 1.31 × 10⁶ Da) was procured from Chem Total labs, Jodhpur, India and Guar Gum (MW 1.41 × 10⁶ Da) from CDH, India. All analytical grade chemicals were used directly to prepare gels in the distilled water.

Methods

Polymer gel preparation

Linear polymer gel samples were prepared by the adding desired quantity of polymer (KG, Guar or CMHPG) in distilled water at 1, 2 and 3 wt.% concentrations with continuous stirring using a magnetic stirrer at 200 rpm for 75 min at 50 °C for improved dispersion and swelling. For the preparation of polymer nanocomposite, first nanoparticles were dispersed in distilled water at desired concentration using solvent based exfoliation via sonication method with high shear and high impact mixing of SiO₂ nanoparticles in solvent (water) for 1 h to improve interfacial bonding between the components and to prevent nanoparticles aggregation (Henrique et al. 2009). Upon completion of nanoparticles dispersion, polymer was added to this solution (Chauhan et al. 2017).

Measurements

Infrared spectroscopy

The infrared spectrum of the KG samples was observed in 3800–400 cm⁻¹ frequency range using a PerkinElmer Fourier transform infrared spectroscopy (FTIR) instrument as described in our previous work, Chauhan et al. 2017.

Morphological analysis

Field Emission-Scanning Electron Microscopy (FE-SEM) and Atomic Force Microscopy (AFM) analyses were conducted on KG and KG–SiO₂ nanoparticles samples using Carl Zeiss FE-SEM model-Supra 55 and Dimension Icon (Bruker AXS, Santa Barbara, CA, USA instruments), respectively, as described in detail in our previous work, Chauhan et al. 2017. The samples were diluted enough to describe the shape and size and also to reduce the aggregation risk upon drying.

Rheological studies

The rheological properties, both static and dynamic, were examined on a Bohlin Gemini rheometer as discussed in our previous work, Chauhan et al. 2017. The prepared gels were examined for thermal stability by observing the viscosity with time at various temperatures (120 °C, 150 °C
and 175 °C) in a high-pressure cell maintained at 900psi (restricted due to the instrumental constraint).

**Polymer breaking studies**

Ammonium persulfate (APS), an oxidative breaker, was used to observe the gel breaking for KG, its nanocomposites, Guar and CMHPG gel samples at 1 and 3 wt% concentration. The gels after adding breaker were left for 24 h at 80 °C to observe degradation with respect to time as discussed in our previous work, Chauhan et al. 2017.

**Sandpack regained permeability test**

The sandpack regained permeability test was performed using flooding experiment (Fig. 1) to determine the damage caused to the prepared sandpack by the broken fracturing fluid. The details of the experiment have been explained in our previous work, Chauhan et al. 2019.

**Result and discussions**

**Functional groups in KG and their variation on addition of nanoparticles**

KG, an acidic polysaccharide, is chemically composed of galacturonic acid, sugars rhamnose and galactose (Chauhan et al. 2017). Figure 2 shows the result of FTIR analysis of KG (powder), KG fluid and KG–SiO2 composite. The characteristic bands present in KG (powder) are: 3436 cm\(^{-1}\) for O–H stretching band of the hydroxyl group, 1731 cm\(^{-1}\) for Acetyl group, 2928 cm\(^{-1}\) for C–H in CH\(_2\) asymmetric stretching and 1614 cm\(^{-1}\) for asymmetrical stretching of carboxylate group or bound water molecules. The bio-polymer on hydration has major peaks at 3436 cm\(^{-1}\) and 2060 cm\(^{-1}\) which reveals that bonding has happened between water and polymer molecules. A wide peak at 3436 cm\(^{-1}\) corresponds to the self-association of hydroxyl groups by hydrogen bonding. The peak at 1036 cm\(^{-1}\) representing the starch content in the powdered KG dissolved for the polymer gel system representing breaking of the inter-molecular starch bonds upon dilution in water and followed by the formation of the hydrogen bond at the open sites with more water molecules leading to swelling of polymer.

The reaction for the composition of inorganic–organic composite structure proceeds rapidly with hydrated silicon dioxide. The presence of acids in KG results in mecha-nochemical dehydration due to acid–base reaction when reacted with silanol group of SiO\(_2\) with the formation of KG–SiO\(_2\) composite. According to the FTIR result, the molecules.
bridging bond Si–O–C is formed on the composite surface due to the reaction in Eq. 1.

\[-\text{COOH} + \text{SiOH} \rightarrow \text{Si} – \text{O} – \text{C} + \text{H}_2\text{O} \quad (1)\]

Figure 3 explains the possible reaction mechanism of KG–SiO$_2$ nanocomposite formation. The molecules of water absorbed on SiO$_2$ act as the bonding link between acids present in KG and SiO$_2$. The hydroxyl groups of acids and sugars present in KG react with acidic centers of SiO$_2$, while the –OH group of –COOH of acids forms hydrogen bond with the molecules of absorbed water, which is again bonded with basic centers on SiO$_2$ surface by hydrogen bonds (Watanabe et al. 2002; Abo-Shosha et al. 2008; Kizil et al. 2002). Thus, FTIR analysis confirms the formation of KG–SiO$_2$ nanocomposite via Si–CH$_2$–Si linkage through –CH$_2$ group and Si–O–C association through –OH group.

**Morphological studies**

FE-SEM and AFM are powerful tools for the structural determination of bio-polymer nanocomposites using nanoparticles as reinforcing material. SEM and AFM images of KG fluid and KG–SiO$_2$ nanocomposite fluid are shown in Fig. 4. SEM images show the spherical appearance of polymer matrix with pores (dark areas) which confirms the amorphous nature of KG and reflects the higher water-binding capacity of the polymer (Fig. 4a, b). Sufficient contrast between nanoparticles and the KG matrix clearly shows the dispersion of SiO$_2$ nanoparticles into the matrix. Even distribution of spherical nanoparticles in the dense polymer matrix could be noticed in the SEM images with some dense agglomeration or voids at some places (Fig. 4c). The morphology of the developed gel was further confirmed from the AFM study.

The AFM phase and the 3D height images for the polymer and nanocomposite fluid are presented in Figs. 4d–f. The phase image also confirmed the spherical network-shaped structure of KG with a more bright contrast than dark one and its 3D height image (Fig. 4e and f) which suggests the involvement of intra- and inter-molecular aggregation. The stiffer surface with high cross-link density is reflected in bright areas (hard silicon phase) and are uniformly distributed over the dark region which

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**Fig. 3** Possible reaction mechanism for KG–SiO$_2$ nanocomposite formation
corresponds to more viscous area with lower cross-link density (KG) (Isabel et al. 2012; Ulaganathan et al. 2012).

The Root Mean Square (RMS) roughness ($R_q$), the average roughness ($R_a$), the skew ($R_{sk}$) and kurtosis ($R_{ku}$) roughness parameters for the KG and KG–SiO$_2$ nanoparticle samples are presented in Table 1. Table 1 shows the synthesized nanocomposite resulted in increase in mean $R_a$ and a decrease in $R_{sk}$ which suggest an increase in the average

**Table 1** Roughness parameters of KG and KG–SiO$_2$ nanoparticles thin films

| Sample      | $R_a$ | $R_q$ | $R_{sk}$ | $R_{ku}$ |
|-------------|-------|-------|----------|----------|
| KG film     | 13.1  | 19.3  | 1.53     | 12.3     |
| KG–SiO$_2$  | 19.7  | 26.9  | –0.501   | 4.77     |
height of particles/agglomerate and dominance of depressions and valleys and is a criterion for well-defined globular structures with good bearing surface, i.e., high load carrying capacity and highly porous fluid film which could provide better shear and thermal resistance of the nanocomposite (Isabel et al. 2012; Kumar and Rao 2012; Hernández et al. 2007; Ulaganathan et al. 2012).

**Linear steady state rheology**

KG and KG–SiO₂ nanocomposite gel samples were investigated for rheological characterization at 1%, 2% and 3% polymer concentrations and 0.25 and 0.50% SiO₂ concentrations for increasing shear rate (0.01–1000 s⁻¹) and temperature (30–150 °C). The rheological results have been compared with Guar and CMHPG which are the most frequently used polymers in the industry (Fig. 5a and b). It is clear from Fig. 5 that viscosity decreases in the order Guar > CMHPG > KG for all three prepared concentrations, and it relates to reducing molecular weight. KG suspensions are able to maintain comparable viscosities due to the presence of acetyl groups, in particularly galacturonic and glucuronic acid residues which promote interactions among polymeric chains and formation of gel network (De Brito et al. 2005). The viscosity increases as polymer concentration increases, mainly as a result of entanglement of the disordered polymer chains which increases with concentration.

Nanocomposite with 0.25% SiO₂ resulted in higher viscosity similar to CMHPG and 0.5% SiO₂ nanocomposite resulted in viscosities similar to guar at higher temperature ranges. Increasing the concentration of nanoparticles resulted in increased viscosity from 625 to 800cP at 150 °C at 100 s⁻¹ for 3% KG and from 180 to 300cP for 2% KG.

The energetically favorable interaction among nanoparticles clusters and polymer chains develops robust physical associations developing highly stable nanocomposite compared to the only polymer. This enhanced strength is mainly because of the viscous and rigid nature of clusters synergy, where the silane (Si–O–C) covalent bonds support the plastic deformation and reversible physical hydrogen bonds (Si–OH and Si–CH₂–C) provide nanocomposites flexibility and elastomeric nature. This synergy offers localized regions of enhanced strength that decay the growth of crack/hole upon deformation (Yang et al. 2013). The cluster formation can significantly influence the transfer of stress and heat in the network due to their porous structure and extensively effective volume fraction. To rephrase it, nanoparticles and polymer associate to develop an intramicelle “pseudocrosslinking,” i.e., a 3D structure via various linkages and surface charge interactions. This results in enhanced viscosity due to robust dynamic polymeric network, direct morphology and rapid energy burst between rigid nanoparticles and soft polymer (Yang et al. 2013; Cassagnau 2013; Ji and Walz 2013; Crews and Huang 2008). From Fig. 5a and b, it could be observed that all the gels samples showed shear-thinning behavior which reduces friction during pumping and is always considered beneficial in a fracturing operation.

The increase in temperature (40 °C to 150 °C) results in thermal thinning of the gel samples. However, from the results in Fig. 6a and b, it is evident that 1% KG gels could maintain the required 90cP viscosity till 80 °C only which can be seen to improve to 120 °C on the addition of 0.50 wt. % SiO₂ nanoparticles. The required 90 cP at 100 s⁻¹ viscosity comes from the rule of thumb applicable for fracturing fluids to suspend proppants and maintain required rheological properties during fracturing and prevailing reservoir temperature (Dantas et al. 2003; Baruah et al. 2014). Contrarily, 2 wt. % KG and its nanocomposites at 0.25 wt.% and 0.50 wt.% SiO₂ (Fig. 6b) always maintained

![Viscosity variation with increasing shear rate for 1 wt.% polymer gel samples at 30 °C.](image1)

![Viscosity variation with increasing shear rate for 2 and 3 wt% polymer gel samples at 30 °C.](image2)
their viscosities above 200 cP at 150 °C which is superior to the recommended value of 90 cP at 100 s⁻¹.

**Oscillatory sweep measurements:**

Figure 7 shows the relationship between the frequency (0.001 to 15 Hz) and the storage modulus \(G'\) and loss modulus \(G''\) at 100 °C for 1 and 2 wt.% linear KG gels and their SiO₂ (0.25 and 0.50 wt.%) nanocomposites, and the various rheological parameters calculated for all the samples is presented in Table 2. No crossover point observed in the results in the studied frequency range indicates the weak dependence of \(G'\) and \(G''\) on frequency and suggests the high particle suspension capability and longer relaxation time for the studied gels. The \(G'/G''\) always retained higher value than the \(G''\), depicting gel behavior with viscoelasticity where elasticity is dominant (Patel et al. 1992; Sodeifian et al. 2012). The \(G'/G''\) ratio values for all the synthesized samples showed a rising trend with the increase in the concentrations of nanoparticles and polymer (Table 2).

The prepared 1 wt.% KG gel samples showed the soft nature of the gel, which implies an imbalance among the polymer chains, leading to defects like loose ends and loops in the polymer network (Grillet et al. 2012). However, the addition of nanoparticles into the polymer matrix resulted in enhanced gelling and elastic properties.

It could be observed from the Table 2 that for 2 wt.% KG gels with and without nanoparticles showed the value of \(G'/G''\) greater than 3 which confirms the true nature of the gels. The increase in \(G'\) could be because of inter-particle interactions or the thermodynamic interactions between nanoparticle surface and the surrounding polymer matrix. The viscoelasticity is controlled by the dynamics of the slick-slip motion between the nanoparticle surface and the polymer chains, which increases the residence time of the entanglement near particle surface and results in reduced mobility of adsorbed chains and significantly diminished frequency dependence as compared to only polymer gel. This implies the formation of pseudo-solid-like three-dimensional

### Table 2 Rheological Parameters for 1 and 2% KG and their nanocomposite at 100 °C in the frequency range 0.01 to 1 Hz

| Sample          | \(G'\) [Pa] | \(G'/G''\) ± SD |
|-----------------|-------------|-----------------|
| 1% KG           | 45.01       | 1.978 ± 0.472   |
| 1% KG–0.25% SiO₂| 72.3        | 2.458 ± 0.233   |
| 1% KG–0.50% SiO₂| 86.3        | 2.896 ± 0.568   |
| 2% KG           | 38.152      | 5.453 ± 2.65    |
| 2% KG–0.25% SiO₂| 41,208      | 5.6789 ± 1.365  |
| 2% KG–0.50% SiO₂| 46,320      | 5.862 ± 1.12    |
Polymer breaking study

Post fracture, gel breaking is necessary so that low viscous broken fracturing fluid can flow back to the surface along with oil and gas. The final viscosity of all the gels broken using APS at 1 and 3% concentration was found to be less than 2cP at room temperature. Table 3 shows the residue generated from Guar, CMHPG, KG and KG–SiO2 nanocomposite, and it can be seen that KG gels yielded least residues. Even KG–SiO2 composite gave the almost similar results as that of KG. The residue generated depends on the amount of natural residues present in polysaccharide, the molecular weight of polymer and the unbroken gel. KG has a lower molecular weight as compared to Guar and CMHPG. The addition of SiO2 nanoparticles in the composite provides significantly fewer residues as the polymer concentration is not increased, and SiO2 does not result in residue formation (Chauhan et al. 2017).

Simulated field conditions rheology

In actual, fracturing fluid during a hydraulic fracturing job is frequently exposed to the HTHP conditions, and it comes in contact with water-sensitive clays. Therefore, it is of utmost significance to study the combined effect of pressure and temperature on the rheological properties under the simulated field conditions, though it has been observed that the rheological properties of fracturing fluid are more sensitive to temperature than the operating pressure owing to its low isothermal compressibility (Fuhua et al. 2012; Sheng-ying et al. 2008). In the present probe, ethanol and KCl have been added to the prepared fluids as temperature stabilizers and an anti-clay swelling agent, respectively. They are frequently added in fracturing fluids to avoid clay swelling and for high-temperature applications. The effect of controlled pressure of 900 psi (restricted due to instrumental constraint) and high-temperature (upto 175 °C) was studied on KG samples and its nanocomposites for one hour under time sweep experiments at 100 s\(^{-1}\) shear rate.

From the results presented in Fig. 8a, it is evident that 1 wt.% KG is able to maintain its viscosity much higher than 90 cP at 100 s\(^{-1}\) at 120 °C and 900 psi which is the minimum required viscosity for fracturing operation (Thampi et al. 2014). The thermal stability was found to enhance on the
inclusion of alcohol and nanoparticles. The 2 wt.% KG and its nanocomposites were found suitable for higher temperature applications above 120 °C as they maintained sufficient viscosity (Fig. 8a, c) up to 175 °C at 900 psi. The inclusion of alcohols increases the polymer solubility in the less polar aqueous–alcoholic solvent system and enhances the polymer swelling (Balaghi et al. 2011; Chauhan et al. 2017).

The applied pressure did not affect the apparent viscosity over the studied time variable, and the viscosity values were found to be sufficient for the 1 h duration showing its effectiveness for fracturing application. The compressibility effect predominant over the applied pressure condition enhanced the thermal stability of the fluid under observation (Luo et al. 2014; Baruah et al. 2016).

**Sand pack regained permeability**

The oil and gas industry faces major challenges using polymer-based fracturing fluid to control the damage caused to the proppant pack and formation because of partial polymeric degradation and insoluble polymeric residues. The water-soluble polymers leave a significant amount of insoluble residues upon breaking, clogging pores, and thereby reducing the effective fracture permeability. The reduction in effective fracture permeability can be studied on a simulated sand pack system as described in our previous work (Chauhan et al. 2019). Table 4 presents the percentage regained permeability of the sand pack system studied by flowing broken fracturing fluids through the sand pack. From the table, it is evident that the linear KG gels and their nanocomposites offered superior regained permeability than guar and CMHPG. The higher regained permeability for KG samples is directly related to smaller molecular weight, size of KG and its better interaction with the breaker as compared to higher molecular weight polymers like Guar and CMHPG. The partial breaking of guar leads to pore blockages and reduced effective permeability (Cooke 1975; Almond 1982; Parker et al. 1994).

**Conclusions**

The KG–SiO₂ nanocomposites offered improved mechanical and rheological properties due to development of active paramagnetic acid and basic centers on the hydrated SiO₂ surface for mechanochemical reactions. The prepared gels were more viscous, elastic and thermally stable as compared to pure polymer gels and resulted in viscosities similar to CMHPG and Guar but leave lower insoluble residues upon breaking. Also, the polymer nanocomposites resulted in increased elastic modulus value and the long time relaxation due to “strong associated particle” effect which leads to their superior proppant suspension capacity compared to all other polymer gels. Therefore, KG–SiO₂ nanocomposite gels could be considered as a promising candidate for their application in the oil industry as fracturing fluid owing to its improved rheology, thermal stability, breaking characteristics and cheaper cost as compared to CMHPG.

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**Declarations**

**Conflict of interest** The authors declare that they have no conflict of interest.

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