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Sustainable production of raw materials from waste cooking oils†

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The current industrial process for recycling Waste Cooking Oils (WCOs) into vegetable lubricants relies on basic decantation and filtration methods, lacking the scientific foundation needed for technical optimization and sustainability. This research addresses these limitations by thoroughly evaluating the technical and environmental impacts of bentonite treatment and water washing techniques. Using a Design of Experiments (DoE) approach coupled with multivariate statistical analysis, key process parameters—temperature, bentonite content, pH, and oil-to-water ratio—were optimized to improve performance and sustainability. Results showed that bentonite had a negligible effect when water treatment was conducted at 75 °C and pH 6 or at 25 °C and pH 2. The efficiency of both recycling methods, as measured by nuclear magnetic resonance spectroscopy and rheological tests, was comparable. However, the green metrics (mass yield, mass productivity, E-factor, and process mass intensity), along with the EcoScale penalty ranking, indicated that water treatment at 75 °C and pH 6 offers the most viable solution. Linear regression of the data acquired through the multivariate analysis driven by the DoE approach provided a mathematical equation which relates the temperature, time, and the oil/water ratio to the equivalent of CO₂ eventually produced. This tool provides recycling industries with a practical framework for optimizing process conditions, balancing technical efficiency with minimized environmental impact, a crucial factor for compliance with green certification programs. The results present a scalable, scientifically validated pathway for the WCO recycling industry to enhance both operational performance and sustainability.

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Sustainability spotlight

It is well established that Waste Cooking Oils (WCOs) represent a valuable raw material for various industrial applications. Outside of the biodiesel industry, the sustainability associated with their use in fine chemicals production has never been thoroughly assessed. In this context, an operationally simple and low-impact recycling process leading to a raffinate ready for further transformation would facilitate the conversion of the high amount of WCOs produced worldwide. The process presented herein was optimized not only in terms of chemical outcomes, but also with consideration for its impact on sustainability. Green metrics and EcoScale applied to key indicators determined through statistical multivariate analysis allowed for the fine-tuning of the recycling conditions, providing a low-impacting protocol of interest for both industrial and academic purposes.

1. Introduction

In the current socio-economic context of increasing scarcity of resources, developing new circular economy models for designing new processes, as well as for the re-definition of existing ones, is a crucial target.^{1–3} This approach has significant potential impact with political support from initiatives like the European Union (EU) Circular Economy Action Plan, 2020,⁴ or the United States (USA) guidelines “Advancing a Circular Economy to Meet Our Climate, Energy, and Economic Goals” of

the White House.⁵ A key step to achieve circular economy models is using wastes as raw materials.⁶ Waste Cooking Oils (WCOs) are ideal for circular productions. WCOs are classified in the European Waste Catalogue under the code CER 20 01 25 as municipal wastes (Directive 2006/12/EC). The EU produces 1–2.5 million tons of WCOs annually, and globally, production is between 41 and 52 million tons.⁷ Looking at the demand of edible oil, its production was estimated in 230 M tons in 2019 (ref. 8) with an increasing rate of 1.5% yearly until 2027.⁹

Chemically, WCOs are comprised of long-chain (C16–C18) triglycerides partially hydrolysed, resulting in varying free fatty acids (FFAs) content.¹⁰ EU regulations mandate WCO disposal before advanced decomposition, monitored by total polar compound levels not exceeding 25–27 wt%,¹¹ to minimize the formation of potential toxic compounds like acrylamide.^{12,13}

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Early disposal, yields waste with high industrial value. Biodiesel industries convert triglycerides into automotive and aviation fuel *via* transesterification.¹⁴ Triglycerides are used in ecological paintings, asphalt binders, and chain lubricants.^{15–17} Hydrolysed triglyceride products, like glycerol and FFAs, serve as raw materials for numerous applications. Glycerol, a biodiesel byproduct, finds use in cosmetics,⁷ animal feed, and chemical processes.¹⁸ It's also a precursor for pivotal chemical intermediates like epichlorohydrin.¹⁹ FFAs find applications in soap, food, surfactants, and foams.^{20–22} Glycerol and FFAs also contribute to low-melting mixtures, offering eco-friendly solvents with favourable green metrics.²³ Isolating pure glycerol and fatty acids from WCO enhances process sustainability, aligning with circular economy principles.

Among WCO's applications with minimal impacting processing, its use as a lubricant is growing, meeting EU Commission environmental standards.²⁴ The bio-lubricant market, mainly in the EU and USA, is projected to grow by 5.6% in 2024.²⁵ The high value of WCOs may lead to the use of edible vegetable oils for industrial purposes, raising concerns about competition between food and non-food applications.²⁶ In the EU, edible oils can't be industrial ingredients,²⁷ posing challenges when sourcing uncertified WCOs from third countries.²⁸ Regarding this scenario, the attainment of a sustainable and circular approach, viable on an industrial scale, for the facile recycling of WCOs becomes significant.

The chemical processes used for reprocessing and recycling vegetable oils, as well as the physical methods for their purification and large-scale industrial implementation, are well-established. However, when these same processes are examined from the perspective of sustainability, significant and often overlooked gaps in knowledge and practical application remain. This work aims to address these gaps by presenting and analysing the purification, recycling, and industrial utilization of waste vegetable oils as lubricants. Our approach serves as a paradigm for the non-food application of end-of-life food products, offering a novel solution that aligns with sustainability goals while enhancing the value of waste materials.

Preliminary studies showed that water washing under controlled conditions, as well as treatment with clays, are both able to purify the WCOs from many minor organic contaminants, providing a regenerated mixture of triglycerides,²⁹ which can be subjected to many chemical transformations depending on the specific production target.³⁰

Within this framework, the present paper addresses the sustainable production of raw materials from WCOs at different levels. First, the industrial feasibility of the proposed recycling process was considered, exploring the best conditions a recycling company should consider customizing its production. To shed some light on the economical, chemical, and environmental sustainability of different possible scenarios, a Design of Experiments (DoE) approach was used to optimize the process of WCO recycling by water washing followed by bentonite processing. Several green metrics³¹ were applied as the atom economy, the environmental factor (E-factor), the environmental quotient (EQ), the effective mass yield, and the mass

intensity.^{32,33} The Van Aken EcoScale³⁴ was finally utilized as a tool for selecting the best production process.

2. Materials and methods

2.1. Chemicals

WCOs were provided by a catering company operating in Milan (Italy). About 100 kg of WCOs produced over six months were collected and used for the present study. Hexane, HCl (37% solution), NaOH, CDCl₃, and D₂O were purchased by Merk Chemicals.

Hydrophilic bentonite was purchased by Merk Europe. Samples of milled bentonites were prepared by ball milling procedure for 20 min in a SPEX 8000 (SPEX CertiPrep, USA) shaker mill operating at 875 rpm.

2.2. Recycling protocol

To remove macro contaminants as food residues, WCO was filtered through metallic sieves. Then, 30 g of WCO were added to 12 (40 wt%) or 3 (10 wt%) mL of water (at pH 2 or 6) and the due amount of bentonite (0 g or 0.6 g). Acidic water solutions were prepared by adding 1 mL of HCl 37% to 99 mL of distilled water. The mixture was then stirred at the selected temperature (r.t. or 70 °C). After 2 h, the resulting heterogeneous mixture was cooled down to r.t. and the phases separated with a separatory funnel. The organic phase was dried on Na₂SO₄, filtered under vacuum, and collected for analysis. The specific conditions of each experiment are reported in the ESI, Table S2.†

2.3. Hydrolysis of the recycled oil

To 100 g of recycled oil (0.33 mol), 3.5 equivalents of NaOH (1.16 mol) dissolved in 20 mL of H₂O were added. The mixture was stirred at 80 °C during 3 h until a solid was formed. The residual water was removed, and the soap washed 2 times with additional 20 mL (2 × 10 mL) of distilled water. HCl (1.2 mol, 120 mL of HCl 37%) was then added to the solid and the mixture stirred for 2 h at 50 °C. Once all the solid part turned into a yellow oil, the mixture was cooled down to room temperature and the organic phase separated from water in a separatory funnel.

2.4. Water recovery

The aqueous phases recovered from the previous described steps were placed into a round bottomed flask and the water distilled under reduced pressure. The residue was then collected for analysis.

2.5. NMR spectroscopy

NMR samples were prepared dissolving *ca.* 25 mg of WCO in 600 µL of CDCl₃ and then transferred to 5 mm NMR tubes. NMR measurements were performed with a Bruker Avance III 400 console (9.4 T) equipped with a direct observe PABBO probehead and a variable-temperature unit (¹H resonance frequency of 400.13 MHz). 1D ¹H spectra were recorded at 302 K with 16 scans using 65 536 points and a relaxation delay of 1 s, over



a spectral width of 20 ppm. Spectra were processed by applying an exponential line broadening ($LB = 0.3$ Hz), and automatically phased and base-line corrected.

2.6. UV-VIS

UV-VIS spectra were acquired with a ONDA SCAN SPECTROPHOTOMETER UV31, in absorbance mode from 1100 to 195 nm, 1.0 step. Few drops of the sample oil dissolved in 1 mL of hexane were placed in a quartz cuvette with 1 cm of path length for analysis.

2.7. Rheology

Rheological tests in steady state rotation were performed with an Anton Paar MCR502 modular compact rheometer, equipped with a Peltier Plate and Hood thermal control systems. Test were carried out at -25 °C, 25 °C, 40 °C and 100 °C, in shear rate control. Shear rate ranged from 1 to 100 s^{-1} . At the temperature of -25 °C tests were also carried out in oscillation, at a constant shear strain amplitude of 0.1%, in the linear regime, and at frequencies ranging from 100 rad s^{-1} to 10 rad s^{-1} . For all experiments a cone and plate (diameter 50 mm, angle 1° , truncation 99 mm) measuring system was adopted.

3. Results and discussion

3.1. Waste vegetable oil (WCO) characterization

To set a starting point in determining the effectiveness of the recycling process, the chemical profile the WCO was characterized by ^1H NMR and UV-VIS spectroscopy (Fig. 1 and 2, Table 1).

The ^1H NMR spectrum of the starting WCO in CDCl_3 is reported in Fig. 1. It is well known that NMR spectroscopy can be used to characterise vegetable oils and several specific techniques, based on ^{13}C , ^{31}P , ^1H and other nuclei, were developed so far (Di Pietro 2020). According to Popescu and coworkers,³⁵ the relative distribution between fatty acids (FAs, *i.e.* linoleic, linolenic, oleic acid) and saturated fatty acids (SFA, as palmitic and/or stearic acid) can be calculated from the ^1H NMR spectrum. In Table 1, the chemical composition of the herein considered WCO is reported and compared with other published samples of similar origin and determined with the same methodology.

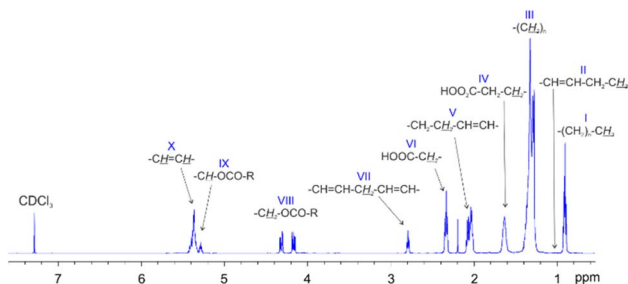


Fig. 1 ^1H NMR spectrum of WCO, with assignment of the characteristic signals.

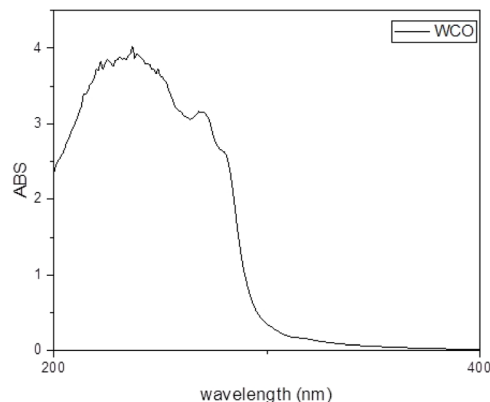


Fig. 2 Portion of UV-VIS spectrum of WCO.

Table 1 Comparison between the chemical composition of different waste vegetable oils (WCO)

Sample	Oleic (%)	Linoleic + linolenic (%)	SFA ^b (%)	IV ^c
WCO ^a	53.3	31.8	14.9	97.0
WCO1 (ref. 36)	66.5	15.9	17.6	81.9
WCO2 (ref. 37)	62.2	20.8	15.4	91
WCO3 (ref. 38)	70	16	16	83

^a Present work. ^b SFA: saturated fatty acids. ^c IV: iodine number.

According to the original methodology of Popescu, the integral corresponding to the signal at 1.02 ppm (II) – assigned to $-\text{CH}_3$ of linolenyl groups – is used to calculate the relative percentages of linoleic and linolenic acids. As it is possible to see in Fig. 1, the intensity of signal II is sensibly lower than the signals I and III, representing the amount of $-\text{CH}_3$ groups of non-linolenyl type and the $(\text{CH}_2)_n$ of the acyl chains, respectively. Thus, the integral of II is not reliable as it partially overlaps to I and III. For this reason, the linoleic and linolenic acids relative percentages are expressed as a sum.

Looking at Table 1, a consistent degree of variability between different batches of WCOs is observed. All the examples reported in the literature are found to be enriched in oleic acid and contain minor and variable amounts of linoleic and linolenic acids. In the WCO herein considered, the total percentage of linoleic and linolenic acid (respectively containing two and three $\text{C}=\text{C}$ bonds) reaches 30%, providing a raw material sensitive to oxidation.³⁹ The iodine number (IV), an empirical parameter useful to define the total amount of unsaturation, shows a value of 97 which lays in the middle between classical edible sunflower vegetable oils (around 100–110), and WCOs (80–90).

Looking at the UV-VIS spectrum reported in Fig. 2, the typical absorption bands of sunflower oil is observed, according with its wide use in southern Europe. From 400 to 1200 nm no relevant bands are detected, discarding the possibility of the presence of EVO oil (extra virgin olive oil) which exhibits typical polyphenolic signals in that region. Indeed, a small band at



310–320 nm, usually associated with flavonoids or with electron transfer processes in triglycerides (ester moiety) is observed. Also, the bands at 240 nm are symptomatic of linoleic acid and referred to the HOMO–LUMO transition of the conjugated dienes.⁴⁰ The UV-VIS profile discards the presence in relevant amount of any organic molecule visible with UV-VIS spectroscopy, confirming the assumption that WCOs usually show low degrees of contamination.

3.2. Process designing

After characterizing the raw material, a recycling process involving water washing and treatment with natural clays was undertaken. Previous studies have shown that these two steps are effective in removing metal particles, small organic molecules, and pigments from waste vegetable oils, thereby enhancing numerous physical properties of the recycled material.^{41–43}

In fact, both recycling processes (water washing and bentonite processing) were reported as effective alternatives and applied to batches of WCOs. However, what is not clear from the published data is which process performs better, especially considering the possibility of a tandem treatment. While these processes were assessed under different conditions, the overall process has never been fully considered or optimized. It is known that bentonite removes small organic molecules, pigments, and even trace metals from vegetable oils, while water washing under controlled conditions partially removes minor organic chemicals, although this aspect has not been explored in detail. To approach the issue, a full factorial 2^k Design of Experiments (DoE) was used to assess the effect of pH, temperature, oil–water ratio (washing step), and clays' amount, on the chemical composition, in terms of oleic acid% and saturated fatty acids%, of the recycled oil (ESI Table S3†).

According to the DoE, a total number of 16 experiments were conducted. The ANalysis Of the Variance (ANOVA) revealed small changes in the chemical composition of the treated oil depending on the condition used as expressed by the Pareto charts in Fig. 3.

Pareto charts show that none of the considered parameters reach the blue line which indicates a P -value < 0.05 . This threshold is highly relevant as discriminates within parameters which stand out as statistically relevant with an interval of confidence $>95\%$. This means that concerning the chemical composition of the oil, expressed in terms of oleic and saturated fatty acids amount, no relevant variations related to the parameters considered are observed. Thus, it is possible to rule-out decomposition pathways during the recycling, independently of the temperature, pH, presence of bentonite. On this basis, a first approach to the recycling optimization was conducted through Surface Response Analysis (SRA). The two statistical models referred to the oleic acid% and to the SFAs% were combined to generate a desirability function (Mannu 2020b). In our case it is desirable increasing one important parameter in lubricants: the oxidative stability. This means having the higher content of oleic acid with respect to linoleic and linolenic acids (one C=C bond *versus* two and three C=C

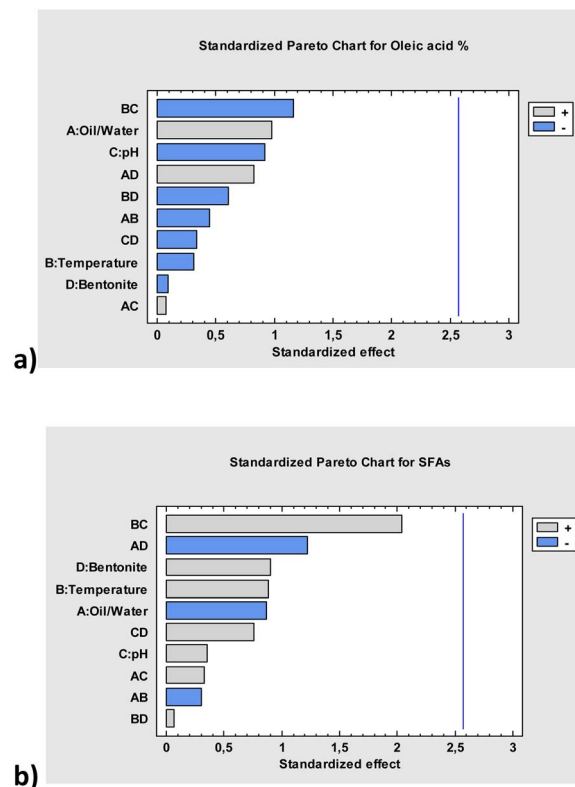


Fig. 3 Pareto chart for oleic acid% (a) and SFA% (b). Blue line marks the P -value < 0.05 .

bonds, respectively). SRA plot shows how the best compromise can be achieved by varying the parameters (Fig. 4).

According to Fig. 4a, the best desirability is achieved without bentonite and an oil/water ratio of 10%. Fixing these two

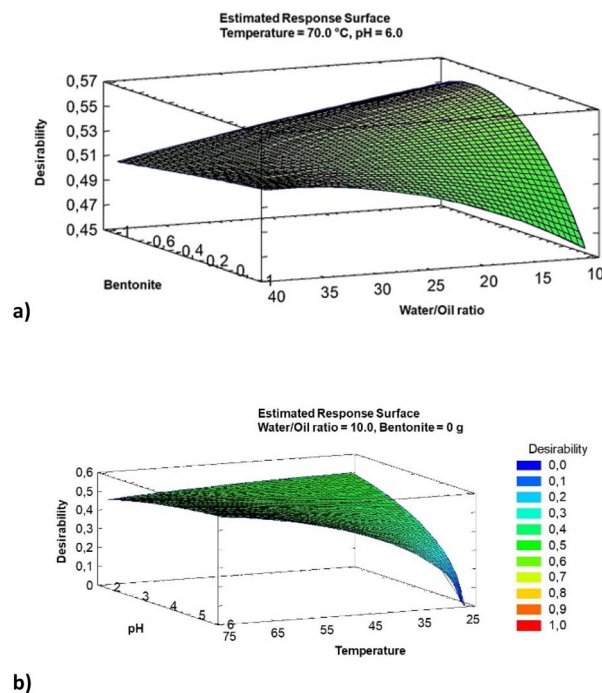


Fig. 4 Surface responding analysis (SRA) for oleic acid% and SFA%.



parameters (Fig. 4b), a relevant effect of the pH combined with temperature is observed. Thus, it is possible to process the WCO at 25 °C if the pH is lower than 4. Alternatively, WCO can be recycled at 75 °C and pH = 6, providing the same outcome. These two alternative options deserve attention as far as the sustainability of the process is concerned. Naively, higher temperatures are detrimental in terms of energy demand. However, acidic pH reduces the possibility to form emulsions during the washing, allowing us to reduce the energy consumption but introduces environmental issues related to the acid concentration.

The information extracted from the statistical analysis represented in the SRA plots is quite impressive. Purification of WCOs can be achieved solely through water washing (at 75 °C, pH = 6, or at room temperature, pH = 2), avoiding the need for bentonite treatment. Additionally, the amount of water can be kept low, with a water/oil ratio of 10. Under these conditions, the amount of chemicals sensitive to oxidation is reduced, resulting in a highly pure mixture of triglycerides being isolated.

Once resolved the technical issues related to the choice of the best recycling procedures, the sustainability aspect was assessed. At first, the CO₂ equivalents (eCO₂) related to the main steps involved (Fig. 5) were calculated (Table 2).

The three most impacting steps on CO₂ equivalents production were highlighted. For each of them, the electricity needed was estimated and then transformed in CO₂. As conversion factor, an amount of 0.373 kg of CO₂ for each kW consumed was considered (Table 2).⁴⁴

Thus, the data relative to eCO₂ were associated to the 16 experiments reported in Table S2† (Table 3).

By taking as response the log[eCO₂], the model resulted very accurate in explaining the variability in eCO₂ generation, with

a $R^2 = 99.995\%$. The Pareto chart reports the six factors with a p -value < 0.05 (Fig. 6).

It's clear that the use of bentonite and the amount of water have a significant impact on eCO₂ equivalents. The interaction between these parameters is particularly noteworthy (see bar CD, Fig. 6). Temperature has a lesser effect, as do the combinations of temperature with bentonite or water amount. The multiple regression model yielded eqn (1), which describes the relationship between estimated eCO₂ equivalents and the four parameters under consideration.

$$\begin{aligned} \text{eCO}_2 = & 0.436528 - 4.1431 \times 10^{-18} \times \text{pH} + 1.25556 \times 10^{-3} \times \\ & \text{temperature} + 8.43333 \times 10^{-3} \times \text{oil/water} + 0.805833 \times \\ & \text{bentonite} \approx 0.436528 + 1.25556 \times 10^{-3} \times \text{temperature} + 8.43333 \\ & \times 10^{-3} \times \text{oil/water} + 0.805833 \times \text{bentonite}. \end{aligned} \quad (1)$$

According to (1), eCO₂ equivalents at various temperatures using different amounts of bentonite and water can be estimated. The pH's effect on eCO₂ is negligible.

Based on the model's results regarding the variation in chemical composition, minimal electricity consumption was achieved under the same conditions. Employing a low amount of water (10 wt%) for washing can effectively recycle WCOs, potentially eliminating the need for bentonite treatment. The optimal operational pH (either 2 or 6) depends on the chosen temperature, resulting in two possible scenarios that were evaluated in terms of green metrics.

3.4. Sustainability and performance assessment

To design a highly sustainable process of industrial interest, the main recycling steps were evaluated in terms of sustainability. The statistical model for estimating eCO₂, as reported above, was complemented by the EcoScale assessment (Section 3.4.1) and several green metrics (Section 3.4.2).

3.4.1. EcoScale. The EcoScale method allows semi-quantitative analysis of a chemical process by assigning penalty points based on specific process features. This tool is useful for comparing processes and selecting the best operative conditions for WCO treatment.

The ideal EcoScale score is 100 and it is calculated using the following equation.

$$\text{EcoScale} = 100 - \sum \text{individual penalty points} \quad (2)$$

To estimate the penalty points Koen Van Aken (Van Aken 2006) defined six sets of parameters characteristic of each process. The

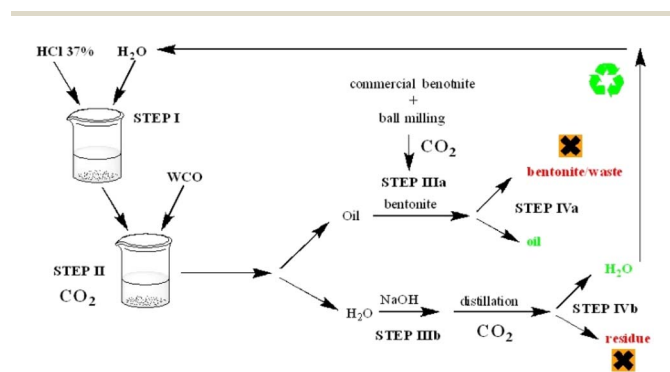


Fig. 5 Scheme of the overall process and eCO₂ impacting steps.

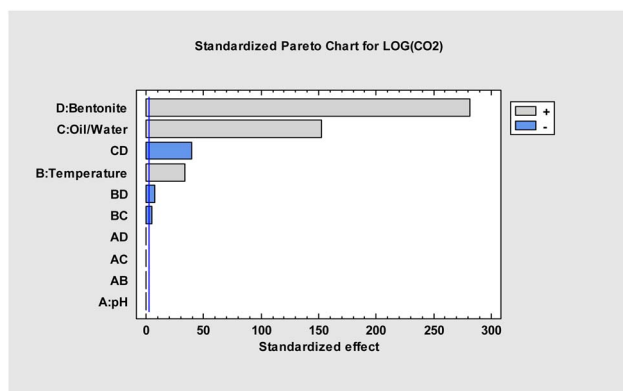
Table 2 Estimated CO₂ produced for each energy-consuming step (ref. 38)

Step	Electric demand (kW)	CO ₂ (kg)	Comments
Water washing and bentonite processing	1.26	0.470	Water washing at 25 °C
	1.42	0.520	Water washing at 25 °C + bentonite processing
	2.52	0.940	Water washing at 70 °C
	2.68	1.000	Water washing at 70 °C + bentonite processing
Water recycling	0.23	0.084	Oil/water = 10%
	0.90	0.337	Oil/water = 40%
Pre-treatment of bentonite	0.03	0.010	For 0.6 g of milled bentonite



Table 3 eCO₂ consumption related to the 16 experiments conducted

Exp.	pH	T (°C)	Water/oil (%)	Bent. (g)	Process (kW)	Milling (kW)	Distillation (kW h)	Total eCO ₂
1	2	25	10	0	0.47	0.00	0.084	0.554
2	6	25	40	0	0.47	0.00	0.337	0.807
3	2	25	40	0	0.47	0.00	0.337	0.807
4	6	70	10	0	0.52	0.00	0.084	0.607
5	2	25	40	0.6	0.94	0.01	0.337	1.287
6	2	70	10	0.6	1.00	0.01	0.084	1.094
7	6	25	40	0.6	0.94	0.01	0.337	1.287
8	2	70	40	0	0.52	0.00	0.337	0.860
9	2	70	10	0	0.52	0.00	0.084	0.607
10	6	70	10	0.6	1.00	0.01	0.084	1.094
11	6	70	40	0	0.52	0.00	0.337	0.860
12	6	25	10	0.6	0.94	0.01	0.084	1.034
13	6	25	10	0	0.47	0.00	0.084	0.554
14	2	70	40	0.6	1.00	0.01	0.337	1.347
15	6	70	40	0.6	1.00	0.01	0.337	1.347
16	2	25	10	0.6	0.94	0.01	0.084	1.034

Fig. 6 Pareto chart for log[eCO₂].

analytic calculations are reported in the ESI (Table S1†). The results relative to the scenario 1 (pH 2 at room temperature) and scenario 2 (pH 6 at 70 °C) are reported in Table 4.

This method was applied to the two optimal scenarios previously defined. The mass yield was similar in both scenarios, but scenario 2 showed a higher score, as HCl is not used. Given the complications of handling toxic and acidic residues, scenario 2

(pH = 6, T = 70 °C) was selected as the optimal WCO treatment process, despite the extra heating required (see Table 5).

3.4.2. Green metrics. Additional green metrics were used to study the sustainability and performance of the process. The mass yield, which determines process efficiency, was calculated by dividing the mass of treated oil obtained by the initial mass of WCO.

$$\text{Mass yield(\%)} = \frac{m_{\text{product}} \times 100}{m_{\text{starting material}}}$$

Mass productivity, a key metric in our analysis, relates the mass of the final product to the mass of all reactants. In our case, it substitutes the concept of Atom Economy as there's no chemical reaction involved. Mass productivity is defined as the mass of refined oil divided by the total mass of raw materials used, including oil, water, and other compounds.

$$\text{Mass productivity(\%)} = \frac{m_{\text{product}} \times 100}{m_{\text{input materials}}}$$

The E factor estimates the total amount of waste generated considering all the process components relative to the final

Table 4 EcoScale for scenarios 1 and 2

	Scenario 1	Penalty	Scenario 2	Penalty
1	Yield: 78.2%	10.88	Yield: 78.7%	10.65
2	WCO (10.7 g, 12.2 mmol)	0	WCO (10.7 g, 12.1 mmol)	0
	Distilled H ₂ O (1.1 g, 61.1 mmol)	0	Distilled H ₂ O (1.1 g, 61.1 mmol)	0
	HCl (37%), (0.4 mg, 0.01 mmol)	0	Na ₂ SO ₄ (99%, 0.8 g, 5.4 mmol)	0
	Na ₂ SO ₄ (99%, 2.4 g, 16.7 mmol)	0		
3	HCl (T, N)	5	None	0
4	Common glassware, stirring	0	Common glassware, stirring	0
5	Room temperature, 2 h	1	70 °C, 2 h	3
6	Density-based phase separation	0	Density-based phase separation	0
	Filtration of Na ₂ SO ₄	0	Filtration of Na ₂ SO ₄	0
	EcoScale score	83.12		86.35



Table 5 Summary of green metrics and EcoScale

Metric	Unit	Scenario 1	Scenario 2	Ideal value
Mass yield	%	78.23	78.70	100
Mass productivity	%	59.22	67.19	100
E-factor	kg kg ⁻¹	0.689	0.488	0
PMI	kg kg ⁻¹	1.689	1.488	1
EcoScale	—	83.12	86.35	100

product mass obtained. Higher E-factor means more waste generated.

$$\text{E-factor (kg kg}^{-1}\text{)} = \frac{m_{\text{total process}}}{m_{\text{product}}}$$

The Process Mass Intensity (PMI) is defined as the ratio of the total input of materials over the product.

$$\text{PMI (kg kg}^{-1}\text{)} = \frac{m_{\text{total process}}}{m_{\text{product}}}$$

Consistent with the EcoScale results, when calculating these metrics for the two previously defined scenarios, the second one exhibits better values than the first in each studied metric (refer to Table 5). Scenario 2 demonstrates higher values of mass yield and mass productivity, indicating greater mass efficiency towards the product. Additionally, the E-factor and PMI are lower, suggesting reduced waste production during treatment. In this context, considering overall feasibility and sustainability, Scenario 2 is evidently more suitable for industrial application (see Table 5).

3.5. Chemical and physical characterization of the raffinate

Following the optimized process (Scenario 2), a first batch of WCO was recycled. Analysis *via* ¹H NMR and UV-VIS confirmed observations made during the optimization study: the main chemical composition of the oil only slightly changes post-recycling. While both ¹H NMR spectra exhibit qualitatively the

same profile for the WCO before and after treatment (see Fig. 7), quantitative analysis of chemical composition reveals an increase in SFA content post-treatment (from 14.9% to 27.5%) at the expense of the unsaturated portion (oleic + linoleic + linolenic percentage decreases from 85.1% to 72.5%). This change is also reflected in the iodine number, which decreases from 97.0 to 82.4, indicating higher oxidative stability.

Comparison of UV-VIS spectra reveal a more resolved portion between 300 and 250 nm. In WCO samples, this region is associated with polyphenols and minor organic compounds, which were removed during the recycling procedure. The band characteristic of the conjugated dienes of linolenic and linoleic acids remains visible and is better resolved at 240 nm (Fig. 8).

3.6 Rheology

Steady state rotational rheometry shows that both WCO and the recycled oil display a Newtonian behaviour (see ESI†) from room to higher temperatures. The viscosities of WCO and recycled oils are very similar and a decrease with temperature according to an Arrhenius dependence, with an energy for the activation of the viscous flow equal to $E_{\text{att}} \approx 167 \text{ J (mol}^{-1} \text{ K}^{-1}\text{)}$ is observed (Fig. 9).

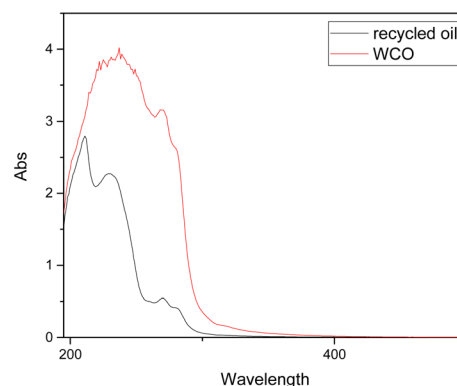


Fig. 8 UV-VIS spectra of the WCO and the raffinate within 200 and 500 nm.

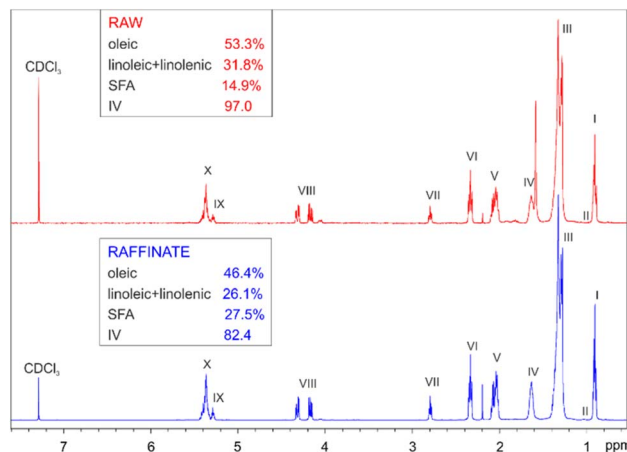


Fig. 7 Comparison between the ¹H NMR spectra of the raw WCO and the raffinate, including the fatty acid profile.

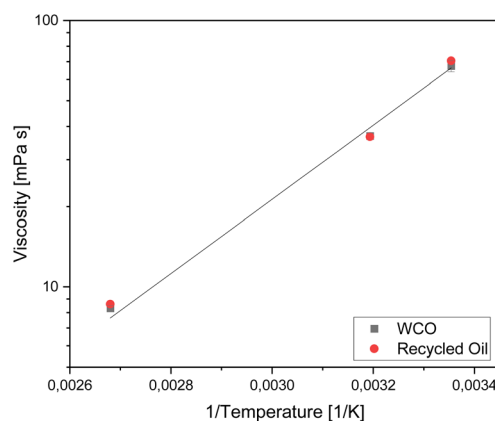


Fig. 9 Temperature effect on the shear viscosity of WCO and recycled oil. The straight line represents an Arrhenius fitting ($R^2 = 0.993$).



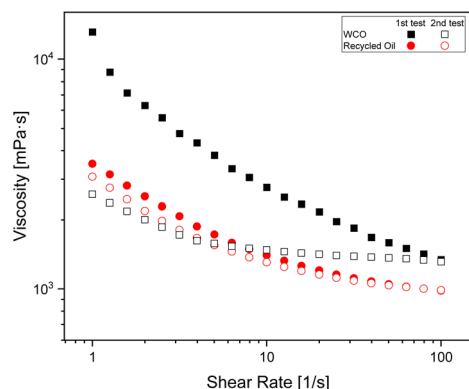


Fig. 10 Viscosity curves of WCO and recycled oil measured in two subsequent flow tests at $-15\text{ }^{\circ}\text{C}$.

Even if no difference can be observed at room and above room temperatures, at sub-zero temperatures some difference can be highlighted between the untreated and treated sample. Indeed, the viscosity curve in Fig. 10 shows a markedly non-linear, shear thinning behaviour; further WCO and recycled oil show significantly different behaviour when subjected to a shear flow in a flow test performed on an “undisturbed sample”, whereas the differences are very limited –or even neglectable–when the sample is tested a second time.

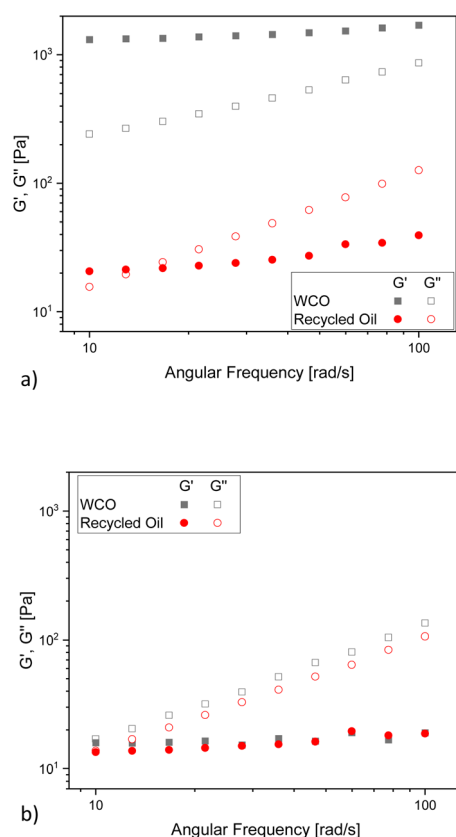


Fig. 11 G' and G'' dependence on angular frequency for WCO and recycled oil (a) before the application of the shear flow in a flow test and (b) after the application of the shear flow in a flow test.

To investigate this feature and its correlation with the sample structure, oscillatory tests were carried out. The G' and G'' curves show a significant difference between WCO and recycled oil before they are subjected to the shear flow (Fig. 11a). Indeed, WCO shows a solid like behavior ($G' > G''$), suggesting the presence of a percolated network for this sample. The shear flow seems to destroy this percolated structure, and the dynamic mechanical analysis performed after flow (Fig. 11b) shows a liquid like behavior ($G' < G''$) comparable for both samples. We speculate that at $-15\text{ }^{\circ}\text{C}$ a crystallization phenomenon starts occurring in both WCO and the recycled oil, but in the case of WCO the presence of contaminants either accelerate the crystallization or contributes directly to the formation of the percolated structure. When a shear flow is applied, this weak network breaks, and does not form again before the second dynamic mechanical test, which makes both material liquid-like and comparable.

From rheology measurements, as well as from UV-VIS data, it is evident that the differences within WCO and recycled oil are small and mostly related with the behaviour at low temperature. This agrees with the fact that the recycling procedure, as proposed, just removes minor chemicals and impurities derived by the life cycle of the oil. Further confirmation to this statement came from the ^1H NMR analysis of the residue obtained by evaporating the washing waters (ESI, Fig. S2†).

3.7. Hydrolysis of the recycled oil

Recycled oil was subjected to alkaline hydrolysis followed by acidification according to Section 2.3 to isolate the corresponding FFAs' mixture. The rate of hydrolysis was monitored by ^1H NMR spectroscopy. In fact, the triplets at 2.3 ppm, related to the CH_2 in a position to the carbonyl moiety, is diagnostic as its chemical shift is very sensible to the chemical environment (ester of acidic function nearby, Fig. 12).⁴⁵

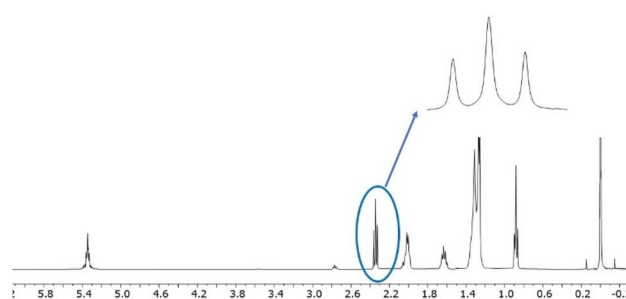


Fig. 12 ^1H NMR (CDCl_3) of FFAs.

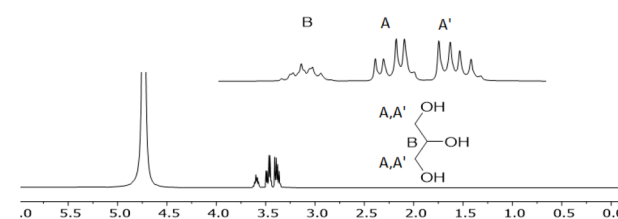
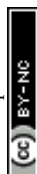


Fig. 13 ^1H NMR (D_2O) of water phase.



On the other side, glycerol was recovered in pure form from the water fraction (Fig. 13).

Conclusions

The use of vegetable oils as lubricants is well-known, with recognized technical advantages such as improved lubricating power, higher flash point, and better adhesion compared to synthetic lubricants. While much research has focused on chemically modified lubricants (*e.g.*, esterified or epoxidized oils), our approach was distinct in utilizing raw triglycerides with minimal transformation, offering a more sustainable alternative to chemically modified waste vegetable oils. Although some industries have begun recycling used vegetable oils through basic purification methods, no optimized or scientifically validated processes were established, leaving a critical gap.

Our work presents a novel contribution by developing a simple yet rigorously optimized process for obtaining clean triglyceride mixtures from used vegetable oils. By employing multivariate statistical analysis, we identified two potential recycling pathways based on chemical composition, specifically the oleic acid percentage and saturated fatty acids (SFAs) content: one at pH 6 and 75 °C, and another at pH 2 and 25 °C—both without the need for bentonite processing. A statistical model capable of estimating CO₂ equivalents (eCO₂) generated by these different scenarios provides an innovative decision-making tool to evaluate the sustainability of recycling options.

The recycling process and the analysis of optimal sustainable conditions were further reinforced by spectroscopic characterization of starting materials, intermediates, products, and impurities (using ¹H NMR and UV-VIS spectroscopy), as well as rheological analysis (steady-state rotational rheometry). Additionally, for alternative applications requiring pure components, we isolated and characterized pure glycerol, and a mixture of free fatty acids obtained through chemical hydrolysis of waste cooking oil (WCO) processed *via* the optimized method.

Through extensive experimentation, we optimized key parameters such as pH, temperature, and water usage, eliminating the need for unnecessary filtration, which is traditionally used in industrial processes. Our method is scalable and adaptable, supported by a mathematical model that allows industries to fine-tune process parameters like temperature, pH, and water-to-oil ratio according to specific operational needs. This approach results in the removal of one unit operation, reducing energy consumption and improving the overall energy balance of the process.

This work aligns with several United Nations Sustainable Development Goals (SDGs), including SDG 9 (industry innovation), SDG 10 (reduced inequalities), and SDG 12 (responsible consumption and production). By converting waste into a valuable industrial resource, especially in regions with abundant waste oil supply, our method promotes circular economy development, with relevance for developing countries. We also provide green metrics and a sustainability ranking to further

validate the environmental, economic, and safety benefits of this approach.

In conclusion, the analytical and theoretical tools developed in this study offer valuable insights into the sustainability and technical feasibility of potential recycling pathways for WCO recovery. The significance of this research, particularly the DoE (Design of Experiments) approach to sustainability assessment, extends beyond the specific case of WCO recycling and serves as a broader framework for environmental research in general.

Data availability

Data are available upon request from the authors.

Author contributions

AM supervision, experimental work, writing and editing, PAF experimental work, writing, MP experimental work, writing and editing, FB experimental work and writing, AnM funding acquisition, supervision, writing-review and editing.

Conflicts of interest

There are no conflicts to declare.

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