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Chitosan-based blended films loaded with cajeput oil as enhancers of antibacterial action against *Staphylococcus aureus* and *Pseudomonas aeruginosa*

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Chitosan-based blended films loaded with cajeput oil as enhancers of antibacterial action against Staphylococcus aureus and Pseudomonas aeruginosa





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Abstract: Chronic wounds (CW) enable pathogen invasion and prosperity, damaging host tissue and impeding tissue remodeling. Essential oils (EOs) exert quick and efficient antimicrobial (AM) action, hindering bacterial resistance. Cajeput oil (CJO) has strong AM properties against *Staphylococcus aureus* and *Pseudomonas aeruginosa* (DOI: 10.3390/antibiotics9060314). Chitosan (CS), a natural cationic polysaccharide, is also AM. CS and poly/vinyl alcohol) (PVA) films (ratio 30/70) were prepared by solvent casting and phase inversion method (DOI: 10.1002/app.48626). Films thermal stability and chemical composition data emphasize polymer blending. Films were loaded with 1 and 10wt% CJO in relation to total polymeric mass. These films reached 0.89 ± 0.05 and 1.14 ± 0.10 mm in thickness, respectively, 23 and 57% thicker than unloaded films. Degree of swelling and porosity also increased, particularly with CJO at 10wt%. CS films showed an outstanding AM activity, eradicating *P. aeruginosa* colony traces within the hour (***p<0.001). Still, loaded CS/PVA films showed improved AM traits, more than unloaded films after 2h of contact (S. aureus: *p<0.05 and **p<0.005 with 1 and 10% CJO, respectively; P. aeruginosa: *p<0.05 with 10% CJO). This study shows that CJO can be dispersed into CS/PVA films and induce bactericidal effects, this way opening new avenues for CW therapeutics.

Keywords: bactericidal; marine-derived polymers; natural bioactive agents; drug delivery systems; blended films.

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Infected wounds

Bacteria are primarily responsible for diabetic foot ulcer (DFU)'s infections, being *S. aureus* the most common bacteria isolated (46.4%), followed by *P. aeruginosa* (22.8%)

S. aureus is a Gram-positive, commensal bacterium



P. aeruginosa is a Gram-negative, invasive bacterium



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The increased resistance of bacteria against antibiotics

serious concerns about DFU therapeutic strategies

Bio-based treatments with quick bactericidal action and low tendency to induce resistance are greatly needed.

Tavares, TD, Antunes, JC et al., Antibiotics 2020, 9(6), 314

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Antibacterial CS



It is suggested that the **antimicrobial activity** of the marine-derived polysaccharide CS results from its cationic nature

Antimicrobial mechanisms

- Electrostatic interaction between positively charged R-NH₃⁺ sites and negatively charged microbial outer cellular components and/or cellular membrane leads to cellular impermeability (inhibiting growth) or cellular lysis (killing bacteria). CS internalization and interaction with cytoplasmic constituents may also occur
- Chelation of metals, suppression of spore elements and binding to essential nutrients to microbial growth interfere with their growth and may contribute to their death

CS's antimicrobial activity is **influenced** by **various intrinsic and extrinsic factors**

CS itself (type, M_{W} , DA, viscosity, solvent and concentration)

environmental conditions (test strain, its physiological state and the bacterial culture medium, pH, temperature, ionic strength, metal ions)



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Antibacterial CJO

Essential oils (EOs):

- aromatic, volatile, lipophilic biomolecules, extracted from regions of plants (e.g. flowers, leaves, twigs, bark, wood, fruits, etc.)
- ✓ formed of complex mixtures of hydrophobic molecules, including thymol, carvacrol and eugenol (among others), which exhibit a broad spectrum of antimicrobial activity against bacteria, fungi, and viruses
- ✓ potential to replace antibiotics due to their inherent and strong antiinflammatory, antiseptic, analgesic, spasmolytic, anesthetic, and antioxidative properties



Chitosan (CS) and Poly (vinyl alcohol) (PVA)



Natural and crystalline polymer Biocompatible and biodegradable Film-forming High viscosity Antibacterial and antifungal properties Ability to absorb exudates

Food and Drug Administration (FDA)-approved as a wound dressing material (topical intended use)



Synthetic and semi-crystalline polymer Biocompatible and biodegradable Film-forming Good mechanical properties: flexibility and swelling capability in aqueous environments

Water-soluble

Multiple FDA-approved medical uses, in the form of transdermal patches, jellies, oral tablets, ophthalmic preparations, intradermal patches and sutures, among others

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- good capacity to form intermolecular hydrogen bonds
- readily forms hydrogen bonds due to a large number of hydroxyl groups

Increase hydrophilicity, improve mechanical properties
Improve stability in aqueous environments

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Main Applications:

Food packaging, controlled release of biomolecules, wound dressing, tissue engineering, membrane bioreactors, pervaporation, reverse osmosis, dye removal, fuel cells

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Solvent Casting + Phase Inversion

CS: 100-300 kDa and 9.6±1.4% DA PVA: 72 kDa and 88% DH



adapted from HP Felgueiras et al., J Appl Polym Sci (2019) doi: 10.1002/app.48626 J. Appl. Polym. Sci. 2018, doi: 10.1002/APF 46188

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Solvent Casting + Phase Inversion



	EO		CS solution		PVA solution		Total 0//\/	\/(m)\	CC(D)/A mass ratios
	m (mg)	V (μL)	m _{cs} (g)	V (mL)	m _{PVA} (g)	V (mL)	- Total %w/V	V _{Total} (mL)	CS/PVA mass ratios
CS	-	-	3.51	39	-	-			100/0
Ρ٧Α	-	-	-	-	3.51	39			0/100
CS/PVA	-	-					9%	39	
CS/PVA/CJO 1%	35.1	39.2	1.053	26	2.457	13			30/70
CS/PVA/CJO 10%	351	392							





Characterization of CS/CJOPVA films



Characterization of CS/CJOPVA films



CS/CLO/PVA film:



Characterization of CS/CJOPVA films



CS/CLO/PVA film:



Antibacterial testing



CS/CLO/PVA film:

S. aureus: the most effective after 6h with 10% EO *P. aeruginosa:* 10% CJO led to an increasingly bactericidal trend,

clear after 2h of contact

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CS film:

S. aureus:

quickest AM action within 1h of incubation

P. aeruginosa: complete bacterial elimination in 1h, effect that endured until tested 24h

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Antibacterial testing



10%

clear after 2h of contact

effect that endured until tested 24h

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Conclusions and Future Work

- CS/PVA blended films were successfully built;
- Both CS and CJO show antibacterial activity against S. aureus and P. aeruginosa;
- ✓ CJO was successfully incorporated in the CS/PVA films at 1 and 10%wt;
- CJO-loaded CS/PVA films were evidently bactericidal effects following 2h of direct contact with the bacteria, being significantly more efficient than unloaded films.
- ✓ Films with 100% CS were particularly more effective than 10% CJO-loaded films against *P. aeruginosa*, by completely eradicating it during the first hour of incubation.

Future work will be directed towards a balance between AM action of CS and its mechanical hindrance after processing, together with the combination with CJO to an intensified antimicrobial profile against both bacteria.

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