## Effectiveness of EDTA Dips on the Shelf-life of Oil Sardine (Sardinella longiceps), Mackerel (Rastrelliger kanagurta) and Prawn (Metapenaeus dobsoni) in Iced Storage\*

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Fresh oil sardine, mackerel and prawn were dipped in 0.1% and 1% solutions of Na<sub>2</sub>EDTA, and stored in ice. Their storage-life was assessed by bacteriological, chemical and sensory methods. Eventhough EDTA treatment controlled the increase in bacterial counts and reduced TMA and TVBN production in oil sardine and mackerel, the consequent beneficial effect was not realised because of the deterioration of fat in these fishes, leading to rancidity. But, for prawn stored in ice, a dip in 1% solution of Na<sub>2</sub>EDTA enhanced the shelf-life by at least 8 days over the untreated control. EDTA absorbed by the muscle of fish and prawn during dip in Na<sub>2</sub>EDTA solution is not completely removed during their iced storage for 25 days.

Earlier works on chemical preservation of fish, particularly from the temperate waters, are well documented by Tarr (1961). Velankar & Kamasastri (1958), Surendran & Iyer (1971 & 1973) and Anand & Setty (1981 a) studied the effect of antibiotics on the bacteria causing spoilage of tropical fish. Possibility of the use of parabens and various other chemicals to preserve fish has been reported by Anand & Shetty (1981 b, c;) and Surendran & Gopakumar (1982 a).

Various sodium salts of ethylene diamine tetra acetic acid (EDTA) have been shown to be effective as dipping solutions for extending the storage life of raw fish (Levin, 1967). Boyd & Southcott (1968) tried Na<sub>2</sub>EDTA and Na<sub>2</sub>CaEDTA for preservation of cohosalmon, leman sole and Pacific cod during refrigerated storage and found that Na<sub>2</sub>Ca-EDTA was not effective, while Na<sub>2</sub>EDTA was. Power *et al* (1968) found that a 1% solution of Na<sub>4</sub>EDTA used as a dip for haddock could extend the storage life in ice for 11 days over the control. However, no work has so far been reported on the application of EDTA to preserve tropical fish, except the studies on the selection of bacterial flora in EDTA treated oil sardine, mackerel and prawn (Surendran&Gopakumar 1982 b). The results of our experiments on the use of EDTA dips to preserve oil sardine, mackerel and prawn in iced storage are presented in this paper.

#### Materials and Methods

Fresh oil sardine, mackerel and prawn were procured from fishing craft operating off Cochin and brought to the laboratory within 2 to 4 h, after catch.

Disodium ethylene diamine tetra acetate  $(Na_2EDTA)$  solutions (aqueous) of 1000 p.p.m. (0.1% w/v) and 10,000 p.p.m. (1% w/v) strength were used as dip solutions. The fish/prawn were dipped in the Na<sub>2</sub>EDTA solutions for 10 min., drained well and packed in ordinary crushed ice, in the fish/prawn to ice ratio of 1:1 and stored in thermocole insulated ice boxes. In all cases, untreated prawn/fish stored in ordinary ice served as the control. The samples were stored for 20–30 days and ice loss was made up by addition of crushed ice, usually on alternate days.

Shelf-life of the fish and prawn was evaluated on the bais of total plate count (TPC), trimethylamine nitrogen content (TMAN), total volatile base nitrogen content (TVBN), volatile acid number (VAN), peroxide value (PV) and sensory qualities. The samples were analysed immediately on the beginning

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of the study and after intervals of 3 to 5 days during storage. TPC was determined using seawater agar (SWA) as described by Surendran (1980). TMAN and TVBN were determined by the Conway microdiffusion method (Conway, 1947) and VAN, PV, moisture and fat by the methods of A.O.A.C. (1960). Sensory evaluation of the samples were made in the raw and cooked state by a taste panel, as detailed by Surendran (1980). The EDTA content of the muscle was determined by the method of Sinclair & Power (1968).

## **Results and Discussion**

# (a) Shelf-life of EDTA treated fish and prawn during iced storage

(1) Oil sardine

Tables 1 and 2 present typical results of the storage study of Na<sub>2</sub>EDTA treated oil sardines. It is evident from Table 1 that the bacterial counts of the control samples increased quite rapidly during storage, while the EDTA treated samples registered lower counts. Of the two concentrations, 0.1 and 1% (w/v), of Na<sub>2</sub>EDTA the sample dipped in 1% Na<sub>2</sub>EDTA had lower counts during the storage. The organoleptic scores of the cooked muscle of the samples showed trends similar to changes in bacterial counts.

The chemical indices like TMAN and TVBN (table 2) showed great variations among the control and treated samples. While the TMAN values of the control sample increased very much during storage, the same in the Na<sub>2</sub>EDTA treated samples remained low, in spite of the fact that such great differences were not reflected in their bacterial populations. However, the changes in VAN and PV values were similar to the bacterial counts as well as organoleptic scores.

### (2) Mackerel

Tables 3 and 4 show the effect of  $Na_2EDTA$  treatment on bacteriological, chemical and organoleptic indices of mackerel held in ordinary ice storage. Only a dip in 1%  $Na_2EDTA$  solution had been tried.

The changes in total bacterial count, TMAN, TVBN, VAN, PV and overall organoleptic scores of the cooked muscle, were more or less similar to those in oil sardines. The TMAN values of the Na<sub>2</sub> EDTA treated fish were appreciably low compared with the control.

Table 1.	Shelf l	ife of	Na <sub>2</sub> EDTA	treated oil	l sardines	stored in ice
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		Concentration	of Na <sub>2</sub> EDTA in	dip solutions
Shelf life index	Days of storage	Control (Na <sub>2</sub> EDTA)	0.1% (1000 p.p.m)	1% (10000 p.p.m)
	0 (before treatment	4.08 x 10 <sup>3</sup> t)	4.08 x 10 <sup>3</sup>	4.08 x 10 <sup>3</sup>
Total plate count/g muscle (TPC)	4 8 12 16 24	7.25 x 10 <sup>4</sup> 2.71 x 10 <sup>6</sup> 5.11 x 10 <sup>7</sup> 3.47 x 10 <sup>8</sup> 2.08 x 10 <sup>9</sup>	1.68 x 10 <sup>3</sup> 5.91 x 10 <sup>4</sup> 2.04 x 10 <sup>5</sup> 2.23 x 10 <sup>6</sup> 1.81 x 10 <sup>8</sup>	2.06 x 10 <sup>3</sup> 3.91 x 10 <sup>3</sup> 1.86 x 10 <sup>4</sup> 9.14 x 10 <sup>4</sup> 3.32 x 10 <sup>6</sup>
Overall organoleptic score of cooked muscle	0 4 8 12 16 24	20 10 6 0	20 12 8 2 0	20 12 10 4 0

Shelf life index	Days of storage	Cont		of Na2EDT. 0.1 (1000 TMAN		1%	, p.p.m) TVBN
	0						
TMAN-mg % and	(before treatment)	1.45	6.40	1.45	6.40	1.45	6.40
TVBN-mg %	4	3.80	14.65	1.45	8.28	1.38	7.25
2 70	8	4.70	18.10	1.60	8.16	1.49	7.68
	12	9.20	24.85	1.75	12.62	1.82	10.55
	16	14.15	29.25	2.20	16.66	1.96	10.76
	24	22.20	36.70	3.86	21.25	2.14	15.28
	0	VAN	PV	VAN	PV	VAN	PV
VAN and	(before	6.72	7.65	6.72	7.65	6.72	7.65
$\mathbb{PV}$	treatment)						
	4	9.86	18.50	7.79	11.28	6.68	6.52
	8	12.75	26.26	8.92	14.65	7.11	10.66
	12	15.08	29.11	11.65	18.22	9.92	12.08
	16	18.65	36.79	14.38	21.67	12.65	18.65
	24	24.28	42.34	21.97	28.08	18.78	24.34

## Table 2. Shelf life of Na2EDTA treated oil sardines stored in ice

 Table 3. Shelf life of Na<sub>2</sub>EDTA treated mackerel during iced storage

Shelf life index	Days of storage in ice	Control (No EDTA)	Dipped in 1% Na <sub>2</sub> EDTA and stored in ice
	0 (before treatment)	2.176 x 10 <sup>4</sup>	2.176 x 10 <sup>4</sup>
Total plate count/muscle	4 7 12 17 24	3.390 x 10 <sup>4</sup> 5.62 x 10 <sup>5</sup> 1.903 x 10 <sup>7</sup> 2.271 x 10 <sup>8</sup> 8.02 x 10 <sup>9</sup>	3.112 x 10 <sup>4</sup> 3.826 x 10 <sup>4</sup> 6.071 x 10 <sup>4</sup> 4.795 x 10 <sup>6</sup> 7.210 x 10 <sup>7</sup>
	TMA		TMAN TVBN
TMAN-mg/100 g.	0 2.	52 7.90	2.52 7.90
and TVBN-mg/100 g muscle	4     3.       7     5.       12     10.       17     15.       24     19.	5019.283825.202528.05	$\begin{array}{cccc} 2.28 & 6.85 \\ 2.70 & 9.42 \\ 3.04 & 12.75 \\ 4.60 & 11.92 \\ 6.75 & 15.90 \end{array}$

## (3) Prawn

Results of a typical storage study of  $Na_2$ -EDTA treated prawns are presented in Tables 5 and 6.  $Na_2$ EDTA solutions of 0.1% and 1% strength have been used for dip treatments.

The total bacterial population of the untreated prawns was always greater compared

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Shelf life index	Days of	Control (No EDTA)		Dipped in 1%	
	storage in ice	VAN	PV	Na <sub>2</sub> EDTA and stored in ice VAN	PV
VAN and PV	0 4 7 12 17 24	5.40 8.98 14.25 20.82 26.65 30.20	6.60 11.85 19.70 26.25 31.75 38.24	5.40 6.20 8.45 9.70 14.24 18.90	6.60 6.78 9.25 14.80 16.22 24.85
Overall organoleptic score of the cooked muscle	0 4 7 12 17 24		20 14 8 4 0 0	20 14 12 8 2 0	

 Table 4. Shelf life of Na<sub>2</sub>EDTA treated mackerel during iced storage

Table 5. Shelf life of Na<sub>2</sub>EDTA treated prawn stored in ice

Shelf life index	Dava of	Concentration	of Na <sub>2</sub> EDTA in	n dip solutions
Shell me mdex	Days of storage Control (No EDTA)		0.1% (1000 p.p.m)	1% (10000 p.p.m)
	0 (before treatment)	2.143 x 10 <sup>4</sup>	2.143 x 10 <sup>4</sup>	2.143 x 10 <sup>4</sup>
Total plate count/g muscle	3 7 12 18 25	6.741 x 10 <sup>3</sup> 4.834 x 10 <sup>4</sup> 2.771 x 10 <sup>6</sup> 1.518 x 10 <sup>7</sup> 1.936 x 10 <sup>9</sup>	7.198 x 10 <sup>2</sup> 3.027 x 10 <sup>3</sup> 1.716 x 10 <sup>5</sup> 1.920 x 10 <sup>6</sup> 6.813 x 10 <sup>8</sup>	6.674 x 10 <sup>2</sup> 1.116 x 10 <sup>3</sup> 5.815 x 10 <sup>4</sup> 6.291 x 10 <sup>6</sup> 3.926 x 10 <sup>8</sup>

with the  $Na_2EDTA$  treated samples. Between the two  $Na_2EDTA$  treated samples, much difference in the bacterial counts was not noticed during the entire iced storage.

The differences in the TMAN values among the control and EDTA treated samples were very significant. While a TMAN value of 22.86 mg/100g muscle of the untreated prawns was reached after 25 days in ice, the corresponding values in the 0.1% and 1% Na<sub>2</sub>EDTA treated prawns were 6.95 mg% and 4.92 mg% respectively.

The TVBN and VAN values exhibited the same trend as that of bacterial counts.

Organoleptically, the Na<sub>2</sub>EDTA treated samples were much superior to the control. The control showed the symptoms of spoilage in 10 to 12 days of iced storage. But the Na<sub>2</sub>EDTA treated samples remained in acceptable conditions even after 16 to 18 days in ice-storage. Even after 18 days of iced storage, the samples dipped in 1% Na<sub>2</sub> EDTA solution did not develop foul smell indicative of spoilage, eventhough the bacteial count exceeded 6 millions per gram muscle.

It can be noted from the results on oil sardine, mackerel and prawn (Tables 1 to 6) that, eventhough the bacterial counts of

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	Concentration of Na <sub>2</sub> EDTA in dip solutions					
Days of storage			0.1 (1000 TMAN	% ) p.p.m) TVBN	1% (1000 TMAN	0 p.p.m) TVBN
0 (Before treatment)	3.26	12.88	3.26	12.88	3.26	12.88
3 7 12 18 25	5.80 7.66 8.15 15.20 22.86	16.20 17.74 19.63 24.14 30.80	3.14 3.70 5.21 5.26 6.95	14.25 16.06 16.28 17.15 21.60	3.40 3.40 4.14 4.75 4.92	12.74 15.80 15.45 17.60 19.85
	VAN	O.S.	VAN	O.S.	VAN	O.S.
0 (Before treatment)	11.60	20	11.60	20	11.60	20
3 ́	14.85	16	11.85	16	11.25	16
•						14 14
		2				14
25	24.15	0	22.05	4	19.88	4
	storage 0 (Before treatment) 3 7 12 18 25 0 (Before treatment) 3 7 12 18 25	$\begin{array}{c} \text{Days of} \\ \text{storage} \\ \end{array} \begin{array}{c} \text{Cont} \\ (\text{No} \\ \text{TMAN} \end{array} \end{array} \\ \begin{array}{c} 0 \\ \text{TMAN} \end{array} \\ \begin{array}{c} 0 \\ 12 \\ 12 \\ 12 \\ 12 \\ 12 \\ 25 \\ 22.86 \end{array} \\ \begin{array}{c} \text{VAN} \end{array} \\ \begin{array}{c} 0 \\ \text{Cefore} \\ \text{treatment} \end{array} \\ \begin{array}{c} 0 \\ \text{TRAN} \end{array} \\ \begin{array}{c} 11.60 \\ \text{(Before} \\ \text{treatment} \end{array} \\ \begin{array}{c} 3 \\ 14.85 \\ 7 \\ 18.08 \\ 12 \\ 17.60 \\ 18 \\ 19.25 \end{array} \end{array}$	$\begin{array}{c c} \mbox{Days of storage} & \begin{tabular}{ccc} \mbox{Control} \\ (No EDTA) \\ \mbox{TMAN} & \begin{tabular}{c} \mbox{EDTA} \\ \mbox{TMAN} & \begin{tabular}{c} \mbox{EDTA} \\ \mbox{TVBN} \\ \end{tabular} $	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

## Table 6. Shelf life of Na<sub>2</sub>EDTA treated prawns stored in ice

the Na<sub>2</sub>EDTA treated samples were almost one log cycle less than the counts of the respective control samples for the same period of storage in ice, the rate of increase in counts did not markedly differ between the untreated and Na<sub>2</sub>EDTA treated samples.

The improvement in shelt-life, as evidenced from TMAN and TVBN values was not reflected in the bacterial counts. Especially the observation that the TMAN values were significantly low in Na<sub>2</sub>EDTA treated fish and prawns in comparison with untreated controls implied that EDTA might preferentially affect TMA producing group of bacteria. Similar observations have been made by Pelroy & Seman (1969) in the case of EDTA treated petrale sole and ocean perch fillets. These authors have reported that a dip in 1% solution of EDTA extended the shelflife at 0.5°C of petrale sole and ocean perch fillets by repressing growth of Pseudomonas spoilage organisms. They found that Na<sub>2</sub> EDTA and Na<sub>4</sub>EDTA were the most effective, extending the shelf-life of ocean perch fillets by 7-10 days whereas Na<sub>2</sub>CaEDTA

gave a 4 day extension of shelf-life over the control fillets.

Power et al (1968) have found that a 1% solution of Na<sub>4</sub>EDTA used as a dip could extend the storage life in ice, of haddock (*Melanogrammus aeglefinus*) fillets by 11 days over that of the untreated controls. They also noticed that TMA production was considerably reduced, but the growth of bacteria was not at all affected by EDTA treatment. Levin (1967), and Boyd & Southcott (1968) have also found that the rate of increase of total bacterial population on haddock and salmon fillets was not affected by Na<sub>4</sub>EDTA.

The fact that uncharacteristic, although not unpleasant tastes were noted in later stages of strorage of EDTA treated fish and prawn suggested that the normal spoilage pattern was altered by EDTA treatment. This idea is further reinforced by the fact (as pointed out earlier) that there was a definite repression of TMA values in the treated samples in comparison with untreated controls, but at the same time, there was no significant difference in total counts. It is possible that EDTA has a selective inhibitory action on the growth of TMA producing bacteria, thus allowing competitive strains to grow at a faster rate (Power *et al.* 1968).

Even though the data presented in Tables 1 to 4 show that EDTA treatments controlled the increase in bacterial counts and reduced the TMAN and TVBN production in oil sardines and mackerel, the consequent beneficial effect was not realised because of the deterioration of fat in these fishes, leading to rancidity, as evident from the rapid increase in VAN and PV values in control and treated samples. Thus EDTA treatment is not useful in extending the shelf-life of fatty fishes. But, for prawns stored in ice, a dip in 1% solution of EDTA enhanced the shelf-life by at least 8 days over the untreated control.

(b) Changes in the EDTA content of the muscle of EDTA treated fish and prawn during ice storage

Concentration of EDTA absorbed by the muscle of oil sardines and prawns during the dip in Na<sub>2</sub>EDTA solutions and the changes in EDTA contents during subsequent storage in ordinary ice are given in Tables 7 and 8.

The quantity of EDTA absorbed by the muscle of both oil sardines and prawns, was directly dependent on the concentration of EDTA in the dip solutions; higher the concentration of EDTA in dip solutions, the greater the amount absorbed.

Table 7. EDTA content of the muscle of oilsardinesdipped in Na2 EDTAsolutionsandsubsequentlystoredin ice

## EDTA concentration micrograms/g muscle

Days of	Dipped in	Dipped in 1%
storage	0.1% Na <sub>2</sub> EDTA	Na <sub>2</sub> EDTÅ
	solution	solution
Just after		
dip	110	326
4	102	290
12	95	246
24	80	212

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Table 8.EDTA content of the muscle of<br/>prawn dipped in Na2EDTA solu-<br/>tions and subsequently stored in ice

# EDTA concentration micrograms/g muscle

Days of storage	Dipped in 0.1% Na <sub>2</sub> EDTA solution	Dipped in 1% Na <sub>2</sub> EDTA solution
Just after dip 3 12 25	196 191 165 157	432 426 384 358

The EDTA content of the muscle of oil sardines, immediately after dip was 110 p.p.m in the case of dipping in 0.1% Na<sub>2</sub>EDTA solution and 326 p.p.m in the case of dipping in 1% Na<sub>2</sub>EDTA solution, the dip time being 10 min. On storage in ice, the EDTA content decreased, but not appreciably. After 24 days of iced storage, the EDTA content of the muscle was 80 p.p.m and 212 p.p.m, for oil sardines dipped in 0.1% Na<sub>2</sub>EDTA solution and 1% Na<sub>2</sub>EDTA solution respectively.

The amount of EDTA absorbed by prawn muscle during dip in Na<sub>2</sub>EDTA solution was greater compared with that absorbed by fish. The EDTA content of prawns dipped in 0.1% Na<sub>2</sub>EDTA solution for 10 min. was 196 p.p.m, which on storage in ice for 25 days, was depleted to 157 p.p.m. Whereas, 432 p.p.m of EDTA was absorbed by prawn muscle dipped in 1% Na<sub>2</sub> EDTA solution for 10 min. and on storage in ice for 25 days, it decreased to 358 p.p.m.

The fact that only a small percentage of EDTA absorbed by the muscle, was washed away during icedstorage might indicate that EDTA was bound or absorbed to the muscle of oil sardine and prawn. That higher amount of EDTA was absorbed by prawn muscle might be due to the fact that, in the case of prawn, the muscle was more exposed and that, owing to the small size of prawns compared with oil sardines, relatively larger areas were exposed to EDTA solution.

Pelroy & Seman (1969) found that the mean EDTA content of ocean perch fillets dipped in 1% solution was 305 p.p.m for the Na<sub>4</sub>EDTA treated fish, 396 p.p.m for the Na<sub>2</sub>EDTA treated fish and 411 p.p.m for the Na<sub>2</sub>CaEDTA treated fish. The residual EDTA contents of haddock fillets treated with 1% Na4EDTA solution were 187 to 333 p.p.m while in haddock fillets treated with 1% Na2CaEDTA solution, the residual EDTA contents varied between 240 and 370 p.p.m. (Power et al 1968). They also found that soft fillets had higher residual EDTA contents than firm fillets of the same size. Boyd & Southcott (1968) have reported that the concentration of Na<sub>2</sub>Ca EDTA in cod samples after immersion for one min. in 0.5% and 1% Na<sub>2</sub>CaEDTA solution was 600 p.p.m and 750-810 p.p.m respectively. The concentration of EDTA in treated samples would undoubtedly be less if fish samples were larger in size. However, the EDTA absorbed by the muscle of oil sardines and prawns, in this study was found to be comparatively less than those reported by the above workers.

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