

The operator submitting the notification completes this form in its entirety. See the guide on how to complete the form for further instructions.

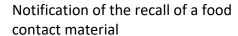
The petification applies to				
The notification applies to	al- fl			
A material or article intended to come into contact with	tn 100d			
Notification has been submitted to				
local food control authority, date and time:				
Name of control authority:		E-mail:		
,				
Finnish Food Authority at takaisinvedot@ruokavirasto	ofi date and time:			
Tillist Food Additiontly at takaismycaotte-faokavirasto	in, date and time.			
2. DETAILS OF THE COMPANY REPONSIBLE FOR T	THE RECALL			
Company name:		Business ID: Telephone:		
Address:		E-mail:		
Type of company operations Manufacturer	Importer			
Contact person regarding recall:		Telephone:		
Address:		E-mail:		
The supplier's contact information in the EU or in a third co	 ountry			
Name: Addre	!SS :	Country:		
,				
3. PRODUCT DETAILS				
Trade name of food contact material:				
General name of food contact material*:				
Toron of front and the state of				
Type of food contact material**:				
		15.1		
Package size or sizes:		Batch code:		
Product description: Attachment Internet li	ink:			
Manufactured by:				
Country of manufacture:	Name of oper	Name of operator who had the product manufactured for it:		
Contact information of the Finnish importer, if different fro	om the data of the re	call company:		
Marketed by:				

^{*}The food contact material's general name and packaging material, serving dishes, tableware, kitchenware, production equipment, etc.

^{**}Type of contact material, e.g. plastic, ceramic, metal, paper and board, etc.



4. DEFECT RESULTING IN RECALL OF THE PRODUCT ANY POTENTIAL CONSUMER COMPLAINTS
Description of defect:
Exceeds the specific migration limit. Which substance migrates?***
The substances that migrate are suspected to be harmful
Contains an unauthorised substance. Which substance?***
Quantity of ingredient used exceeds the specific limit. Which substance?***
Ingredient is suspected to cause colour, taste or odour nuisance in food
Misleading marketing. What claims have been used in marketing?
Incorrect labelling on the packaging (consumer products). Which defect?
Other defect, please state?
How was the defect detected?
Regulatory control RASFF notification In-house control
Information from another operator (e.g. from manufacturer to importer) Consumer or customer complaint
Other reason, please state?
Have complaints about the product been received when the defect was detected or previously?
Yes. When and how many?
□ No
Analysis results (attached). The results must indicate at least the following: who took the samples, what was analysed, the results.
Pending analysis results will be submitted to:
Local food control authority, date:
Finnish Food Authority, date:
*** To be stated if known.
5. DETAILS OF THE SALE AND CIRCULATION OF THE DEFECTIVE PRODUCT
Where have the products been on sale in Finland?****
If the product has been on sale in an online store, who administers the online platform?
For how long the defective product has been on the market?
How much of the defective product has been manufactured/imported?
How much of the defective product is estimated to still be on the market?
now much of the defective product is estimated to still be on the market?
Is the defective batch / Are the defective batches on the market in other EU countries?
Yes, in which country/countries?
No Not known
Is the defective batch / Are the defective batches of the product on the market in third countries?
Yes, in which country/countries?
No Not known
****Please state, for example, the address of the online store or the name of the stores to which the product has been supplied for sale
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6. RECALL ACTION	AND	COMM	1UNICA	TIN
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Recall action taken or planned (who, what, when?)		
Communication done or planned to consumers (newspapers, press releases, etc.)		
What will happen to the recalled products?		
They will be brought to merchantable condition by e.g. re-labelling		
They will be returned to the seller / manufacturer in the country of origin		
They will be disposed of, how?		
Other, please state?		
Other information relating to the product and recall		