

Measuring the electron neutrino mass with bolometer detectors

Monica Sisti

Università degli Studi di Milano-Bicocca

& INFN - Sezione di Milano

on behalf of the Milano, Como and Trento groups



Incontri di Fisica delle Alte Energie 2004

Torino 14-16 aprile 2004

Direct neutrino mass measurement

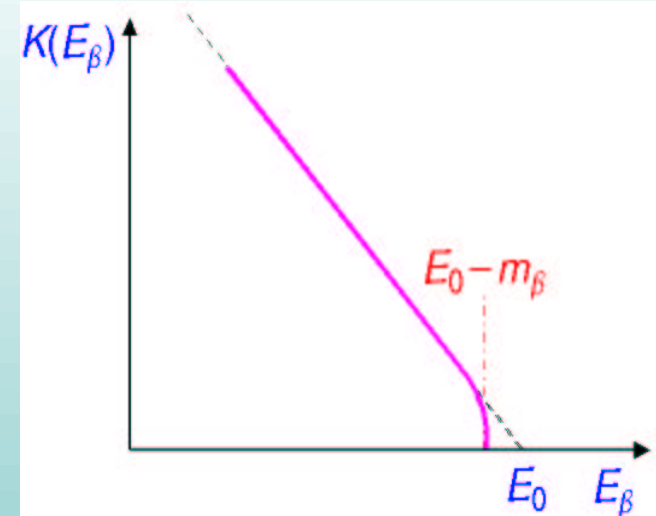
- ◆ neutrino oscillation evidence \Rightarrow finite m_ν
- ◆ neutrino oscillations give only Δm^2
- ◆ to fix absolute scale \Rightarrow study of β decay

kinematics:

- ◇ gives electron antineutrino mass
- ◇ 2 possible experimental approaches
 - ▷ spectrometry (β source \neq detector)
 - ▷ calorimetry (β source \subseteq detector)

calorimeters measure all the energy released in the decay except for that carried away by the $\bar{\nu}_e$

due to the different systematic effects calorimetric and spectrometric approaches are complementary



- in case of mixing with 3 degenerate neutrinos:

$$m_\beta \equiv \frac{\sum m_i |U_{ei}|^2}{\sum |U_{ei}|^2}$$

- in case of no mixing:

$$m_\beta \equiv m_{\bar{\nu}_e}$$

Calorimetric approach /1

- ◆ **Calorimeters measure the entire spectrum at once**
 - ⇒ use low E_0 β decaying isotopes to achieve enough statistics near the end-point
 - ⇒ best choice ^{187}Re : $E_0 = 2.47$ keV
 - ⇒ $F(\delta E = 10 \text{ eV}) \sim 2 (\delta E/E_0)^3 = 1.3 \times 10^{-7}$
- ◆ **Calorimetry advantages (in principle)**
 - ▲ no backscattering
 - ▲ no energy losses in the source
 - ▲ no atomic/molecular final state effects
 - ▲ no solid state excitation
- ◆ **Calorimetry drawbacks**
 - ▼ systematics due to pile-up effects
 - ▼ energy dependent background

Calorimetric approach /2

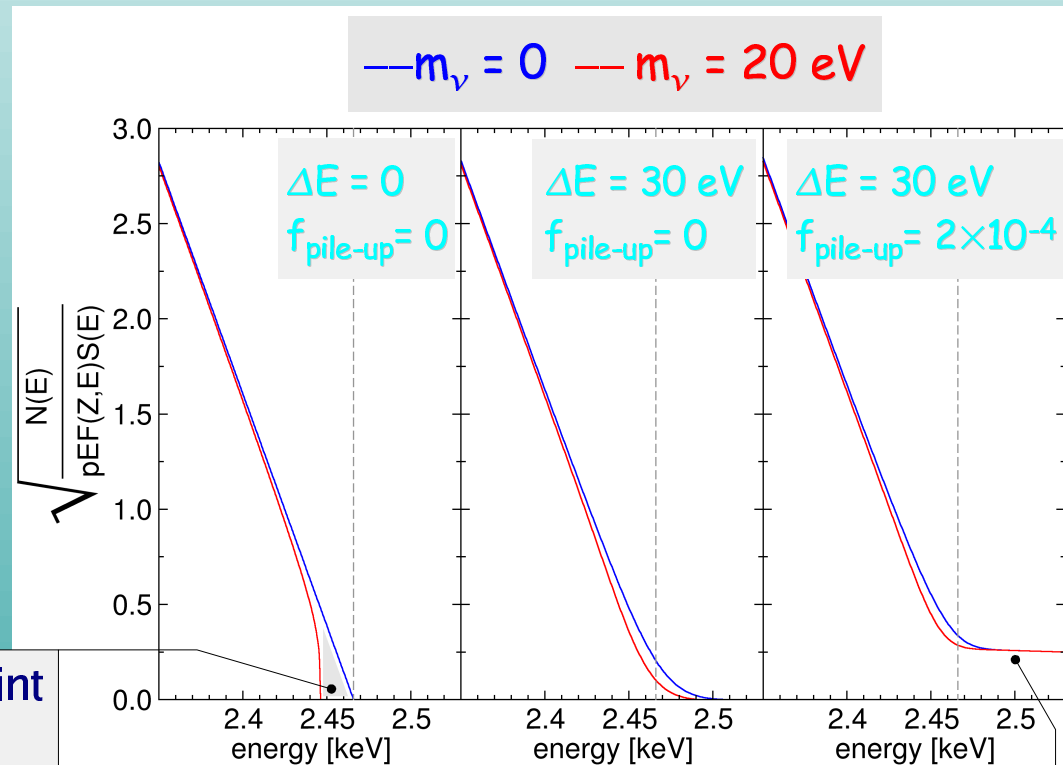
- time unresolved superposition of β decays

Pile-up:

- for a source activity A_β , a time resolution τ_R and an energy resolution function $R(E_\beta)$:

$$N^{\text{exp}}(E_\beta) \approx (N(E_\beta) + \tau_R A_\beta \cdot N(E_\beta) \otimes N(E_\beta)) \otimes R(E_\beta)$$

Pile-up and energy resolution effects:

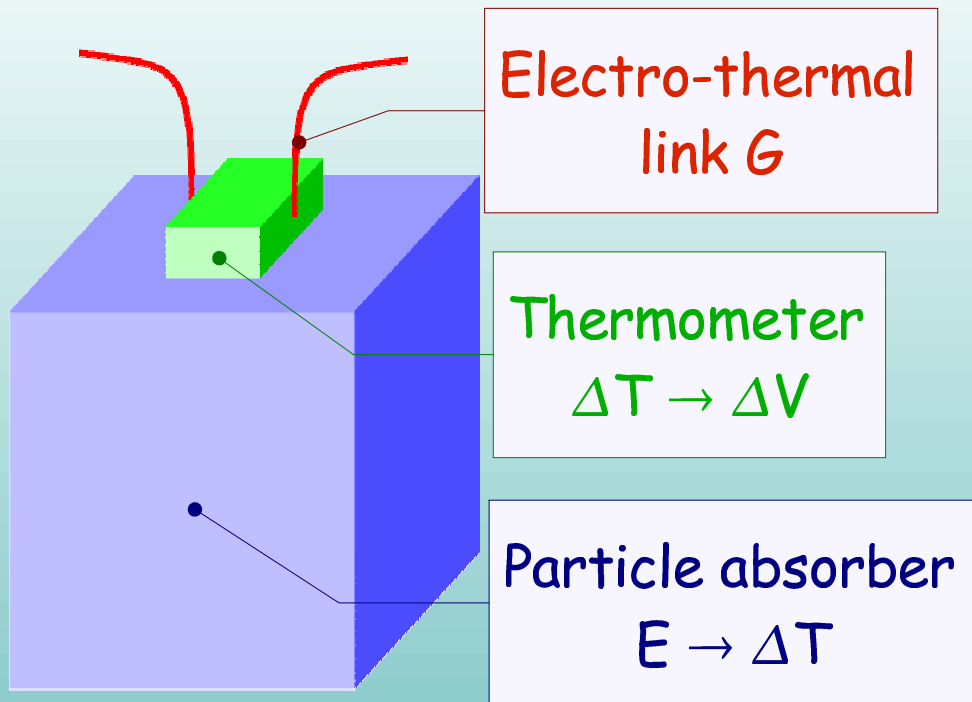


fraction F of decays below the end-point

$$F(\delta E) = \int_{E_0 - \delta E}^{E_0} N(E_\beta, m_{\bar{\nu}_e} = 0) dE \approx 2 \left(\frac{\delta E}{E_0} \right)^3$$

pile-up fraction: $f_{\text{pile-up}} = \tau_R A_\beta$

Thermal detectors



Detection principle:

- $\Delta T = E/C$
 C thermal capacity
⇒ low C
→ low T (i.e. $T \ll 1K$)
→ dielectrics, superconductors
- ultimate limit to energy resolution: statistical fluctuation of internal energy U

$$\langle \Delta U^2 \rangle = k_B T^2 C$$

Thermal detector properties

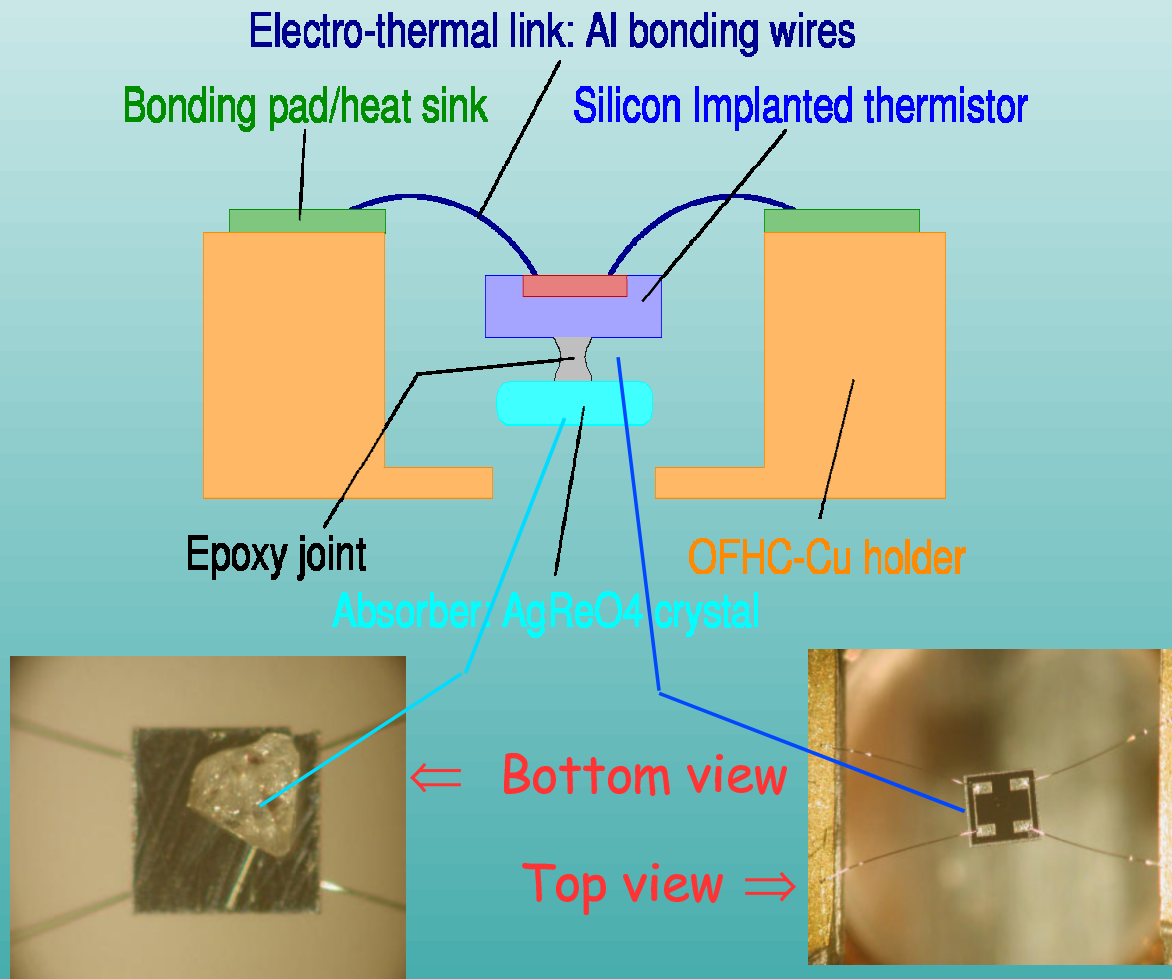
- ▲ high energy resolution
- ▲ large choice of absorber materials
- ▲ true calorimeters
- ▼ slow $\tau = C/G \sim 10^{-1} \div 10^3$ ms

- example: 1 mg of Si @ 100 mK
 $C \sim T^3$ (Debye) $\Rightarrow C \sim 10^{-13}$ J/K
6 keV X-ray $\Rightarrow \Delta T \sim 10$ mK
 $\Rightarrow \Delta U \sim 1$ eV

Milano μ -calorimeters for ^{187}Re β decay study

Neutrino mass measurement with arrays
of 10 AgReO_4 μ -calorimeters.

- ⇒
- lower pile-up
 - higher statistics



Absorbers

AgReO_4 single crystals

^{187}Re fraction ~ 0.32

$A_\beta \approx 5.4 \times 10^{-4} \text{ Hz}/\mu\text{g}$

Mass 250 ~ 300 μg



Measurement statistics and Detector performance

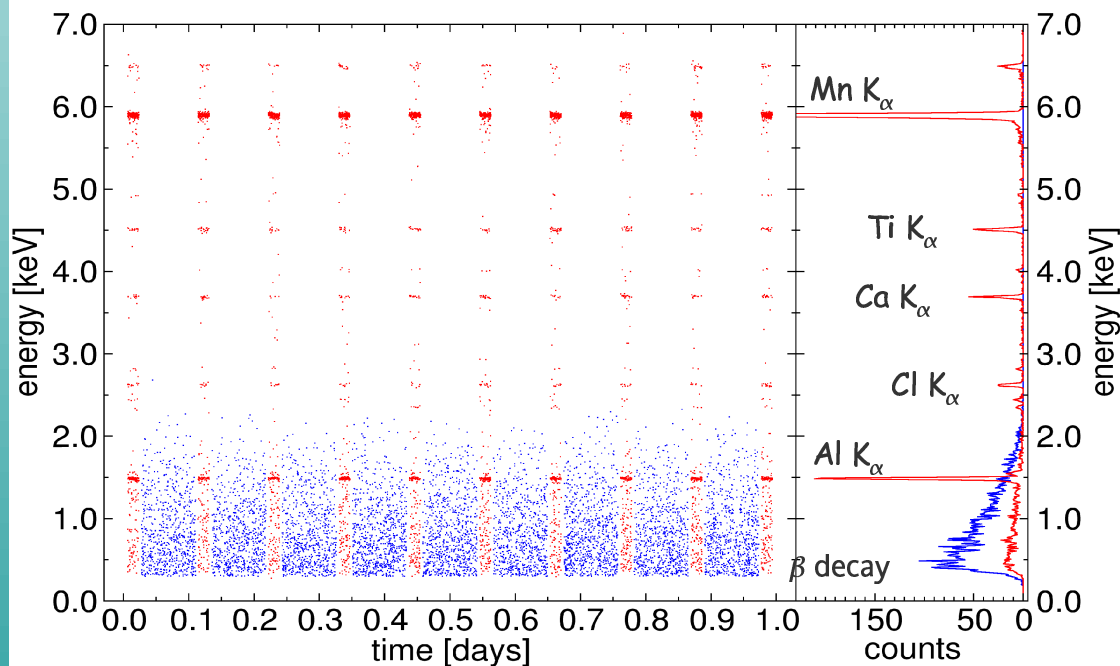
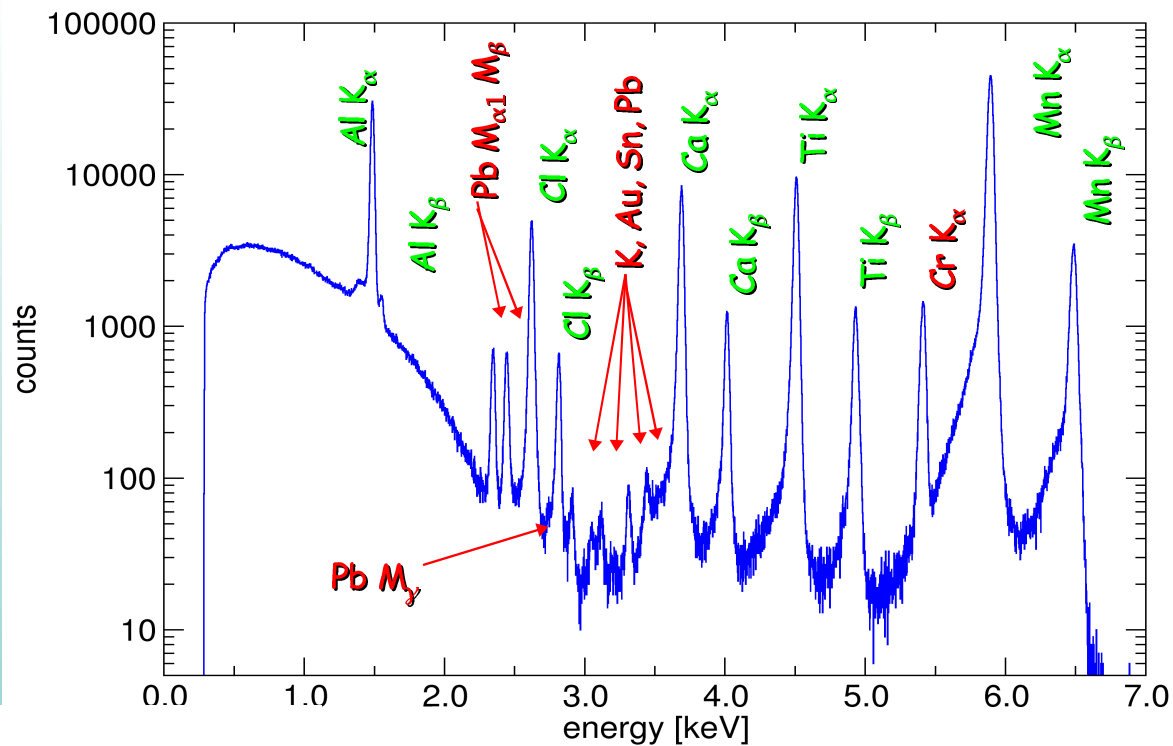
- Measurement started on June 21st 2002 and ended on April 29th 2003
 - ↳ 313 days of measurement (including daily service to cryostat, test calibrations, ...)
- Total acquired AgReO₄ mass: 2.174 mg ⇒ ¹⁸⁷Re activity: 1.17 Hz
- total statistics of β decay acquisition: 168 days ➔ "beta acquisition efficiency": 54%
 - ↳ 8751 hours×mg (AgReO₄)
 - ↳ 16.8×10⁶ ¹⁸⁷Re decays
- total statistics of calibration (X-ray source open): 42 days ↳ 2168 hours×mg (AgReO₄)
- Total effective measuring time: 210 days ➔ "measurement efficiency": 67%

	AgReO ₄ mass [μg]	T _{op} [mK]	R _{op} [MΩ]	shutter ON [h]	shutter OFF [h]	Beta threshold [eV]	ΔE _{FWHM} baseline [eV]	ΔE _{FWHM} at 1.5 keV [eV]	ΔE _{FWHM} at 2.6 keV [eV]	τ _{rise} [μs]
1	272.0±3.0	104.0	1.3	996.7	4037.2	350	20.1	23.7	26.3	440
2	259.0±6.0	91.3	1.9	991.7	4069.2	350	19.4	23.2	26.6	340
3	280.3±6.5	104.0	1.3	1005.9	4105.2	350	25.8	28.7	30.5	670
4	227.7±3.2							bad performances		375
5	249.7±2.0	107.1	1.2	996.3	3803.2	470	21.0	25.4	28.9	590
6	284.0±3.0	86.9	2.2	1006.7	4075.2	350	22.5	25.7	29.2	500
7	281.0±3.7							broken crystal		370
8	282.0±3.4	72.2	4.0	1013.9	4106.4	400	24.6	29.3	33.4	510
9	268.7±5.9	96.7	1.6	981.5	4017.1	400	18.4	21.3	24.9	430
10	278.3±1.5	101.3	1.4	981.9	3967.1	700	23.9	29.2	34.5	445
TOT:				7974.65	32180.7		21.9	25.5	29.1	
							Array summed spectrum			

Calibration spectra

7975 hours × detector with
fluorescence source open

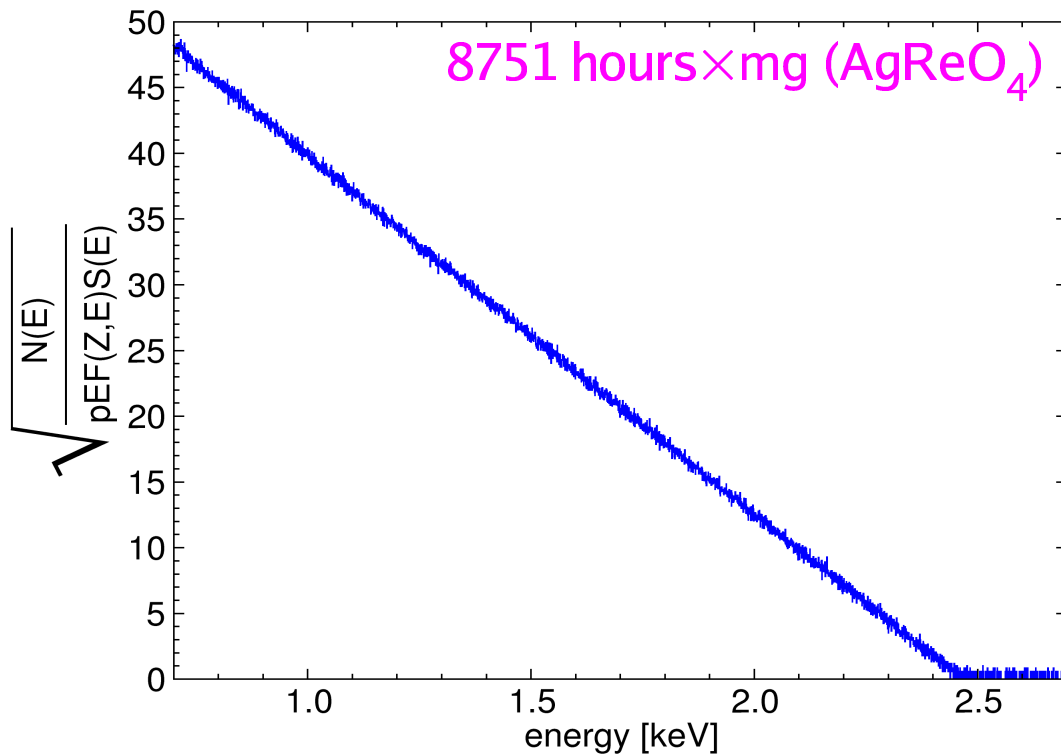
a typical single channel one-day
measurement ↴



- many extra peaks due to fluorescence of surrounding materials
- $Cl\ K_{\alpha}$ position nominal position ($E_{Cl\ K_{\alpha}} = 2.6217\ keV$) missed by 0.3 eV

^{187}Re β spectrum - summed Kurie plot

- 6.2×10^6 ^{187}Re decays collected above 700 eV
- Fit with the function: $F = (f_{\text{th}} + f_{\text{pup}} + f_{\text{bck}}) \otimes f_{\text{det}}$
 - f_{th} → theoretical spectrum
 - $f_{\text{pup}} \propto f_{\text{th}} \otimes f_{\text{th}}$ → pile-up spectrum
 - f_{bck} → background spectrum
 - f_{det} → detector response function



Free fit parameters:

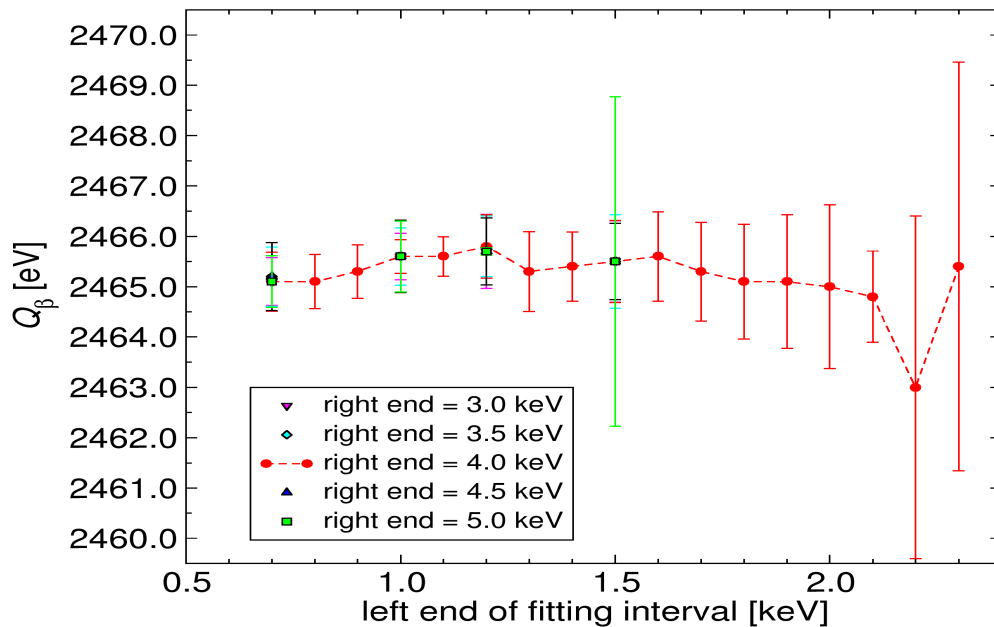
- β end-point
- m_{ν}^2
- β spectrum normalization
- pile-up spectrum normalization
- background level

^{187}Re end-point energy

$$Q = 2465.3 \pm 0.5_{\text{stat}} \pm 1.6_{\text{sys}} \text{ eV}$$

Fitting interval: $0.9 \div 4 \text{ keV}$

→ Most precise value for ^{187}Re



Systematic errors come from uncertainties in:

- ✓ theoretical function for ^{187}Re decay;
- ✓ detector response function and energy resolution;
- ✓ background shape below the β spectrum.

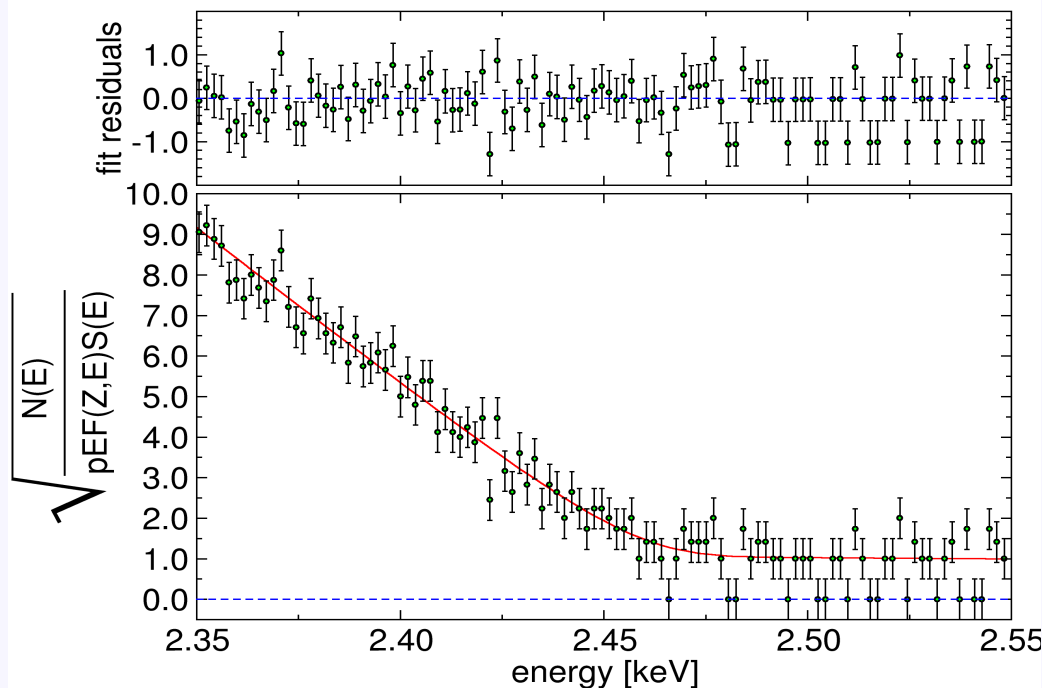
→ Q value is stable with respect to fitting interval \Rightarrow good description by fit function F

Neutrino mass

$$m_\nu^2 = -112 \pm 207_{\text{stat}} \pm 90_{\text{sys}} \text{ eV}^2$$



$$m_\nu < 15 \text{ eV (90 \% C.L.)}$$



Fitting interval: $0.9 \div 4 \text{ keV}$

- ◆ gaussian response function with $\Delta E_{\text{FWHM}} = 28.5 \text{ eV}$
- ◆ free constant backg.: $7 \times 10^{-3} \text{ c/keV/h}$
 - ◇ S/N (Q-30 eV ÷ Q) ~ 12
- ◆ free pile-up fraction $f_{\text{pile-up}}$: 1.9×10^{-4}
 - ◇ $\chi^2/\text{d.o.f.} = 0.905$

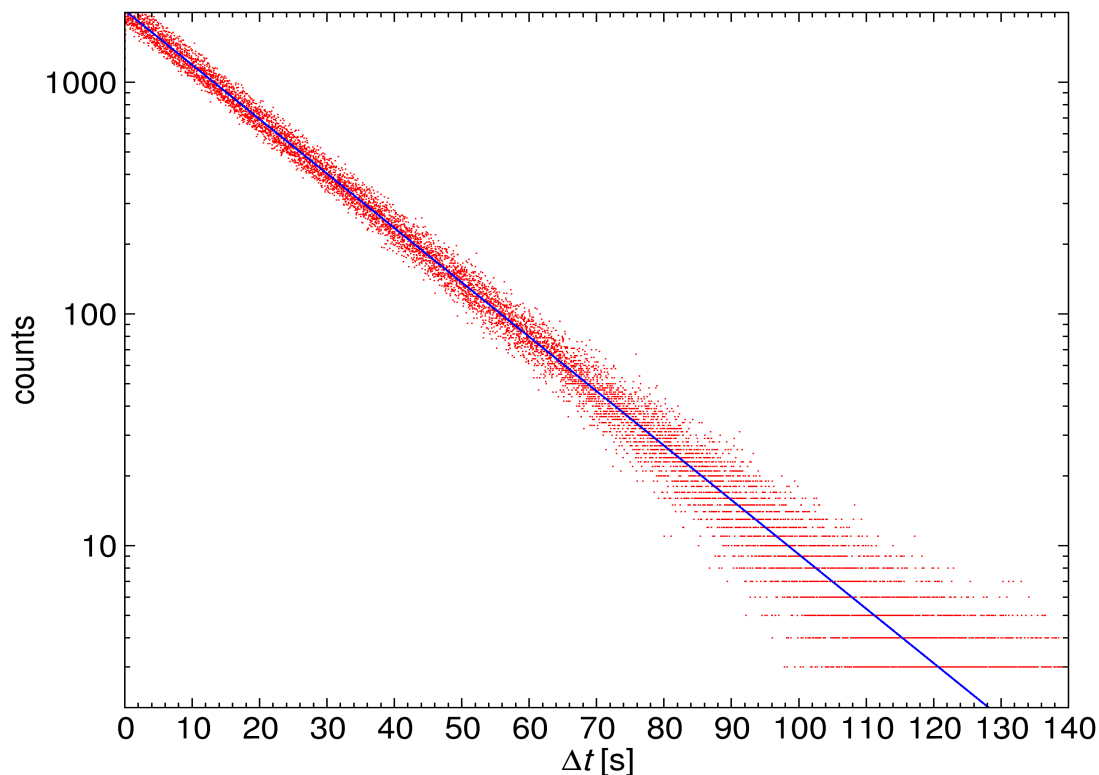
→ same sources of systematics as for the end-point

Preliminary results on a smaller data set in:
C.Arnaboldi et al., Phys. Rev. Letters 91 (2003) 161802

^{187}Re half lifetime

$$\tau_{1/2} = 43.2 \pm 0.2_{\text{stat}} \pm 0.1_{\text{sys}} \text{ Gy}$$

- ◆ from distribution of time intervals Δt between ^{187}Re β decays
- ◆ statistical error mainly from uncertainties in the masses
- ◆ systematic error mainly from uncertainties in the pile-up discrimination
 - ◇ most precise published value; especially relevant for geochronology

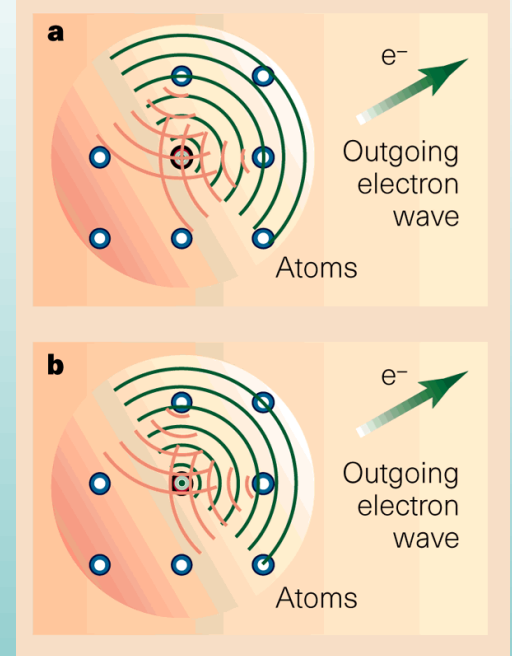


$$N(\Delta t) = A e^{-r \Delta t}$$

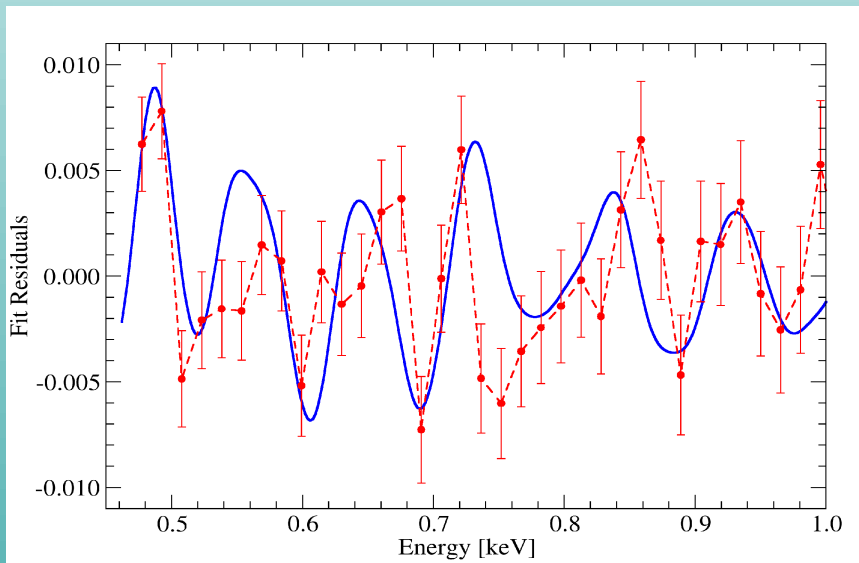
$r \equiv \beta$ decay rate
 $r \propto 1 / \tau_{1/2}$

Beta Environmental Fine Structure (BEFS)

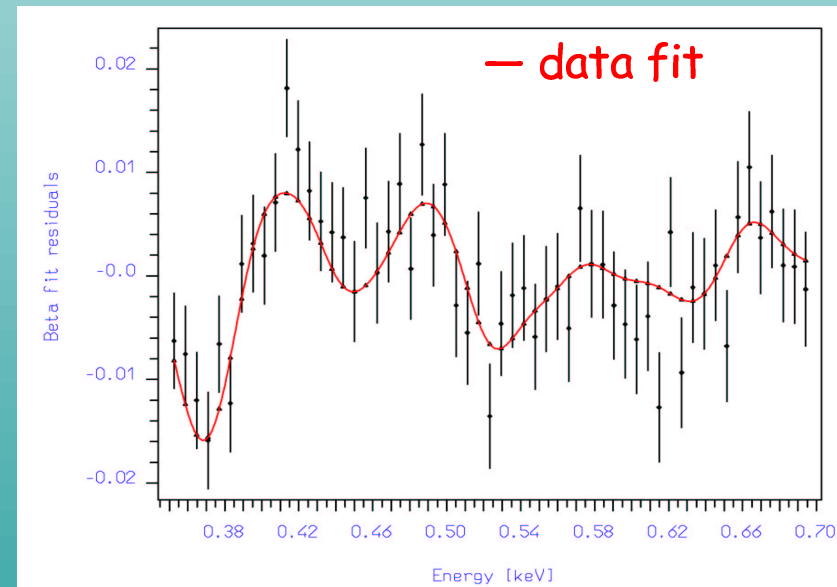
Modulation of the electron emission probability due to the atomic and molecular surrounding of the decaying nucleus: it is explained by the wave structure of the electron



BEFS experimental evidence in AgReO_4



↪ analysis still in progress ...



Future strategy

an experiment for the measurement of m_ν with an approach other than the spectrometric one is important

phase I: experiment that reaches a sensitivity of $\approx 2 \text{ eV}$ before the end of Katrin experiment

phase 2: experiment with a sensitivity similar to that of Katrin $\approx 0.3 \text{ eV}$

	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013	2014
Katrin				0.3 eV							
phase I			2 eV								
phase II (?)						0.3 eV					

phase I is needed:

- because it's the only possible one with present technology
- to check systematics

$$\Sigma(m_{\bar{\nu}_e}) \propto \sqrt[4]{\frac{E_0^3 \Delta E}{A_\beta t_M}} \propto \sqrt[4]{\frac{1}{N_{\text{eventi}}}}$$

m_ν sensitivity estimation by Montecarlo method

Simulation inputs

- ▷ $N_{ev} = N_{det} \times t_M \times A_\beta$ total number of events
 - ▼ N_{det} number of detectors
 - ▼ t_M measuring time
 - ▼ A_β ^{187}Re activity for single detector
- ▷ $f_{pile-up} \approx \Delta t \times A_\beta$ pile-up event fraction
 - ▼ $\Delta t \approx 3\tau_{rise}$ time resolution for pile-up identification
- ▷ $g(E)$: gaussian energy resolution function
 - ▼ ΔE FWHM detector energy resolution

experiment

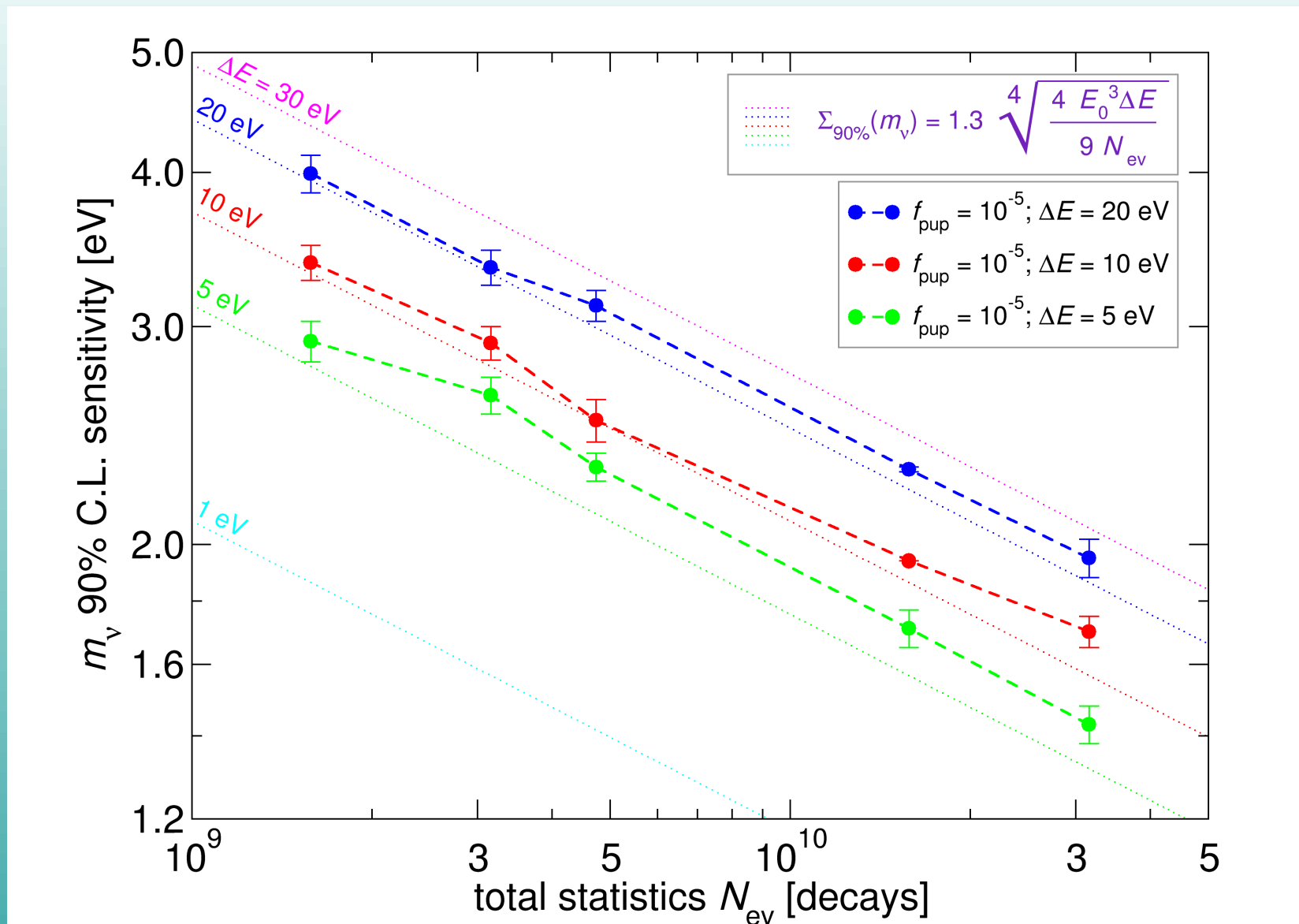
N_{det}	8
t_M [y]	0.59
$\langle A_\beta \rangle$ [dec/s]	0.15
$\langle m_{AgReO_4} \rangle$ [μg]	271
N_{ev} [$\times 10^6$]	16.7
$\langle \tau_{rise} \rangle$ [μs]	490
$\langle \Delta E \rangle$ [eV]	28.5
$\langle b \rangle$ [c/keV/det]	26.3
m_ν 90% CL limit [eV]	15



N_{ev} [$\times 10^6$]	17
$f_{pile-up}$	2×10^{-4}
ΔE [eV]	29
b [c/keV]	210
m_ν 90% CL limit [eV]	16 ± 1

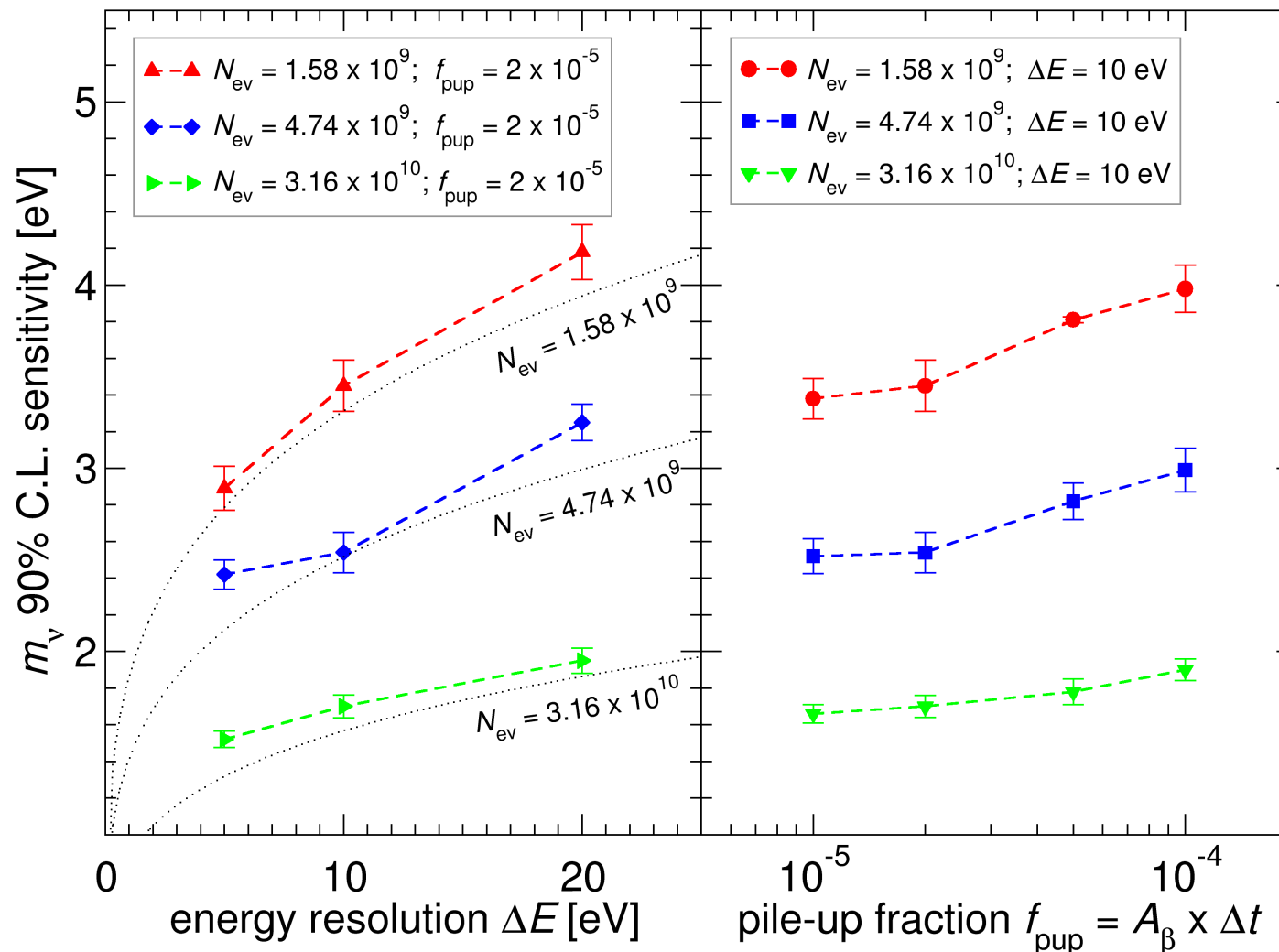
MC simulation

m_ν sensitivity vs. statistics

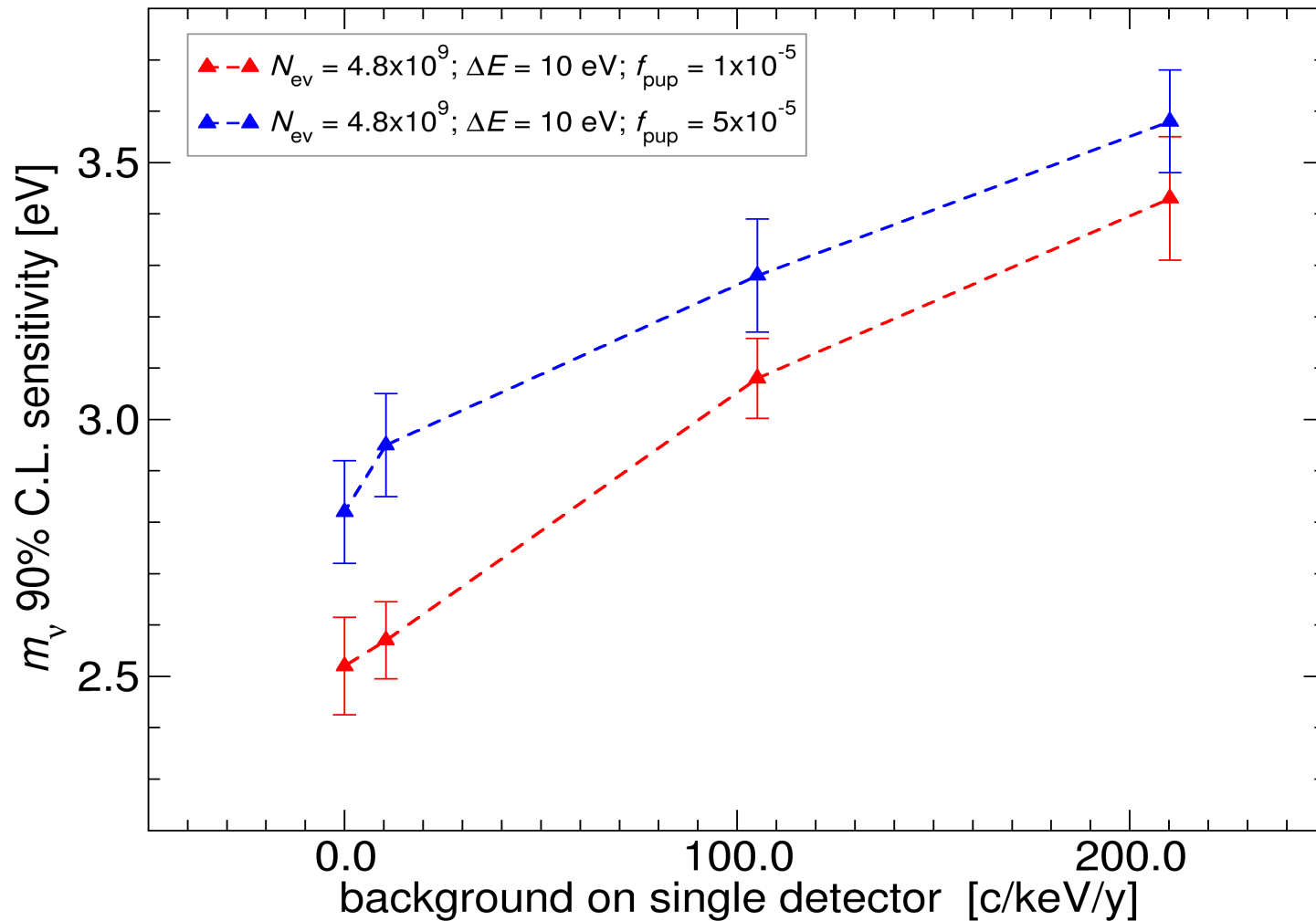


m_ν sensitivity vs. energy resolution

m_ν sensitivity vs. time resolution



m_ν sensitivity vs. background



Phase I: experimental configuration

Montecarlo input parameters			90% CL sensitivity	Possible experimental configurations			
N_{ev} [$\times 10^9$]	$f_{pile-up}$ [$\times 10^{-5}$]	ΔE [eV]	m_ν [eV]	N_{det}	t_M [y]	$\langle A_\beta \rangle$ [dec/s]	$\langle \Delta t \rangle$ [μs]
1.4	2.0	10	3.5	100	2	0.20	100
3.2	2.5	10	3.0	200	2	0.25	100
4.7	2.5	10	2.5	200	3	0.25	100

our present μ calorimeters

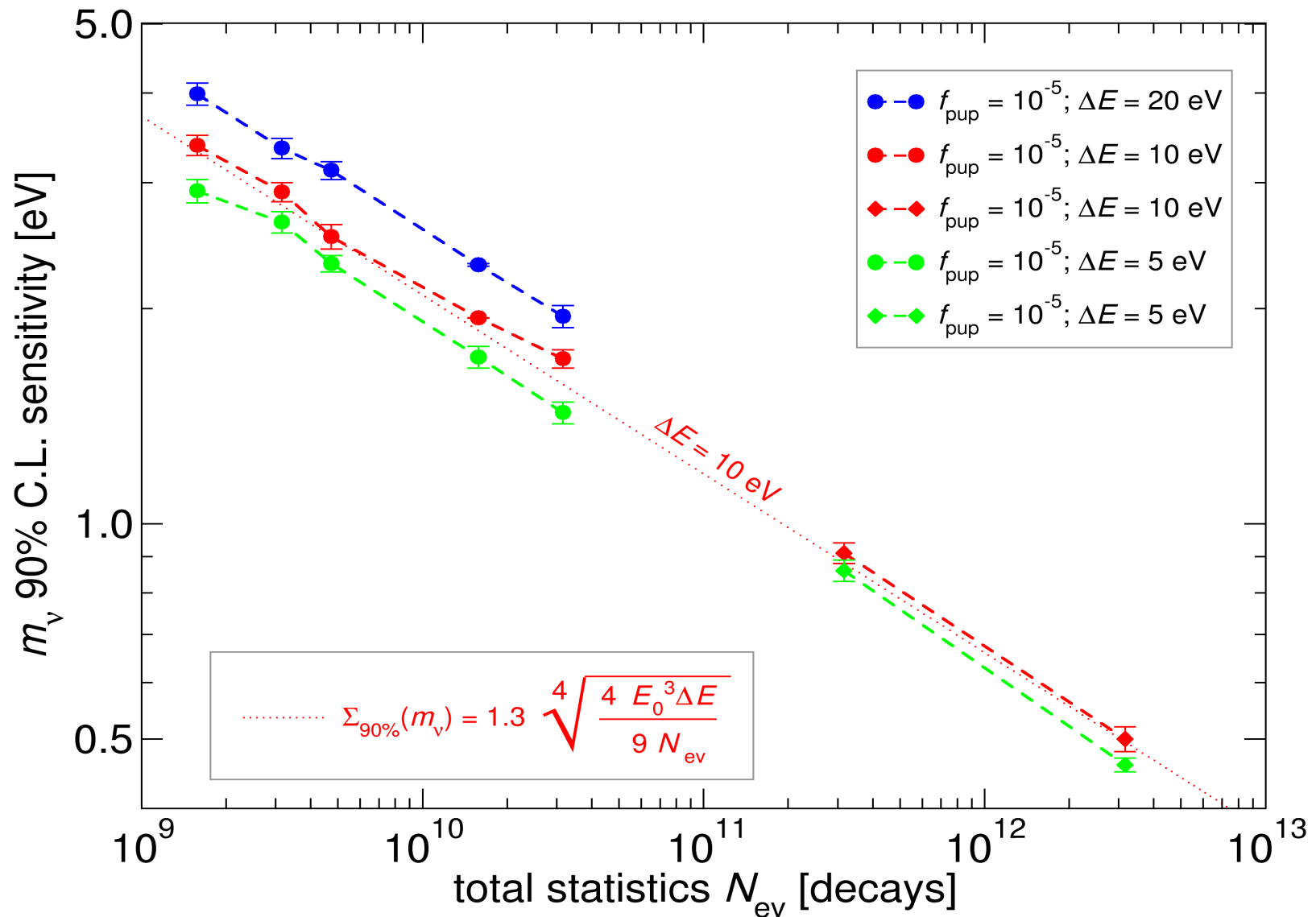
- $m_{AgReO4} = 250 \div 285 \mu g$
 - ▶ $\langle m_{AgReO4} \rangle = 270 \mu g$
 - ▶ $\langle A_\beta \rangle = 0.15 \text{ Hz}$
- $\tau_R = 340 \div 670 \mu s$
 - ▶ $\langle \tau_R \rangle = 490 \mu s$
- $\langle f_{pup} \rangle = \langle \Delta\tau \cdot A_\beta \rangle = 2 \times 10^{-4}$
 - ▶ $\Delta\tau \approx 3 \cdot \tau_R \approx 1 \div 2 \text{ ms}$
- $\Delta E_{FWHM} = 24 \div 34 \text{ eV}$
 - ▶ $\langle \Delta E_{FWHM} \rangle = 28 \text{ eV}$



required μ calorimeters

- $A_\beta = 0.25 \text{ Hz}$
 - ▶ $m_{AgReO4} = 460 \mu g$
- $\langle f_{pup} \rangle = \langle \Delta\tau \cdot A_\beta \rangle = 2.5 \times 10^{-5}$
 - ▶ $\Delta\tau \approx 3 \cdot \tau_R = 100 \mu s$
- $\Delta E_{FWHM} = 10 \text{ eV}$

Phase II: m_ν sensitivity vs. statistics



Phase II: possible experimental configurations

Montecarlo input parameters			90% CL sensitivity	Possible experimental configurations			
N_{ev}	$f_{pile-up}$ [$\times 10^{-4}$]	ΔE [eV]	m_ν [eV]	N_{det}	t_M [y]	$\langle A_\beta \rangle$ [dec/s]	$\langle \Delta t \rangle$ [μs]
3.2×10^{10}	1.0	5	1.7	500	2	1	100
3.2×10^{12}	1.0	5	0.5	1000	10	10	10
3.2×10^{13}	1.0	5	0.3	1000	10	100	1

- **goal: be an alternative to the KATRIN experiment**
- conditions to meet to go for phase II:
 - ▷ phase I must be successful
 - ▼ no unexpected source of systematics
 - ▷ a very large collaboration is needed
 - ▷ really innovative techniques are required
- phase II results would come much later than the KATRIN experiment
 - ▷ it must be also considered that KATRIN could fail

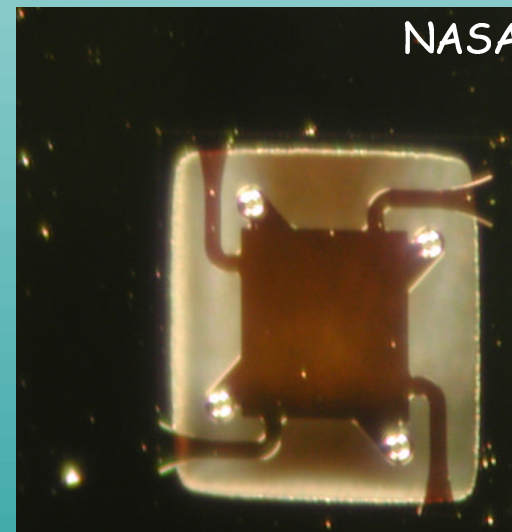
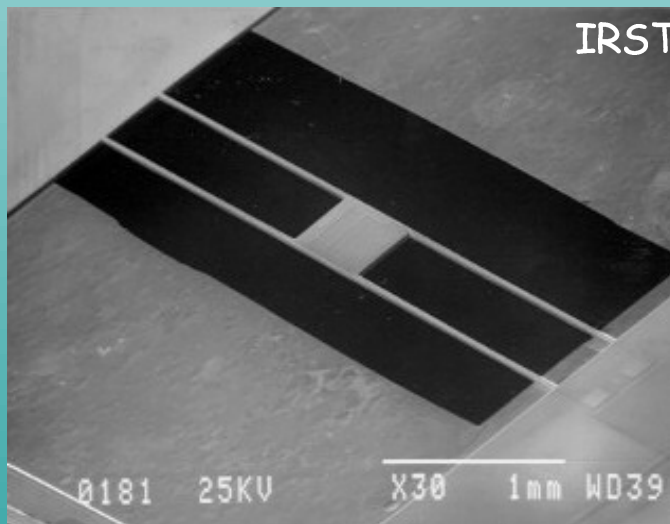
Conclusions

- ✓ Bolometric detectors, even if relatively young, are giving interesting and competitive scientific results (e.g. double beta decay, dark matter, etc.)
- ✓ Calorimetric technique for the direct measurement of the neutrino mass is an appealing complementary approach
- ✓ The Milano neutrino mass experiment with microbolometers has set an upper limit of 15 eV on m_ν after less than 1 year of continuous running
- ✓ A new experiment to improve this limit is now starting the R&D phase

more info at <http://crio.mib.infn.it/wig>

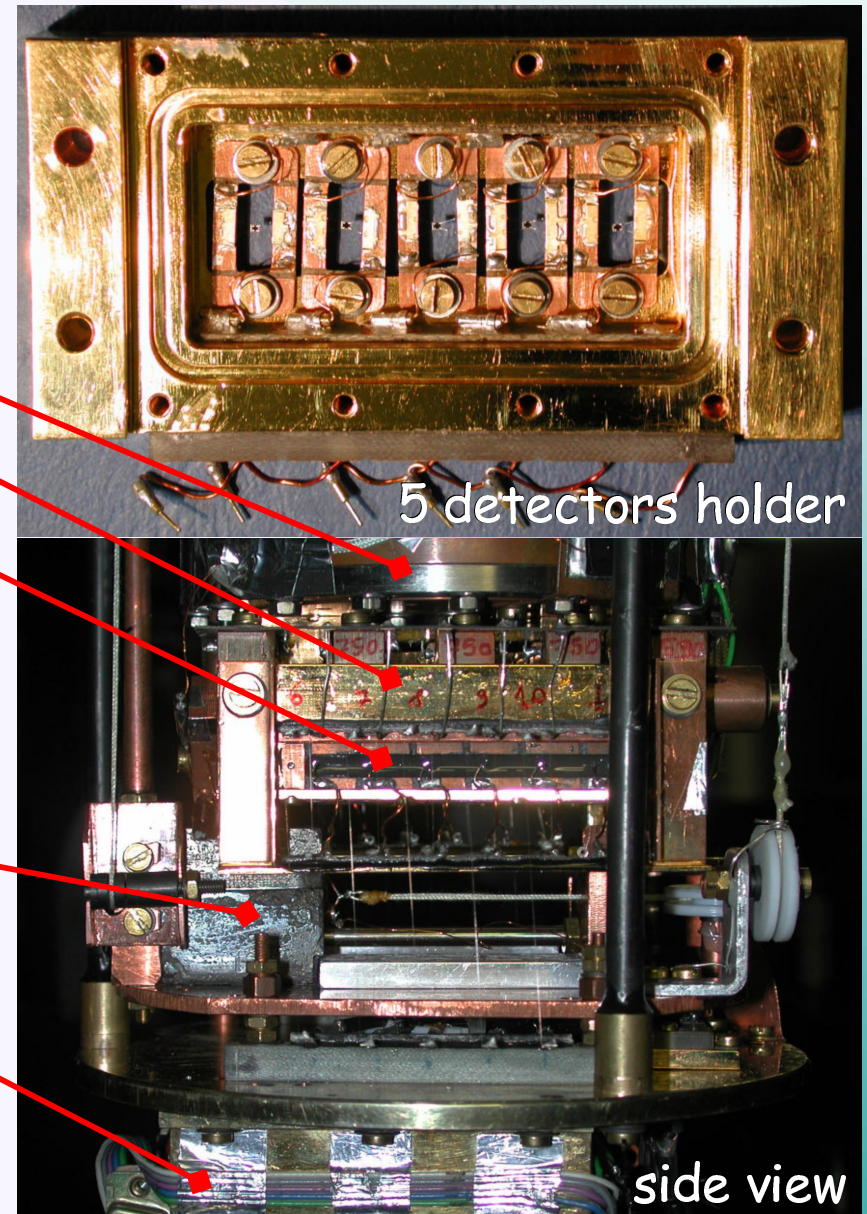
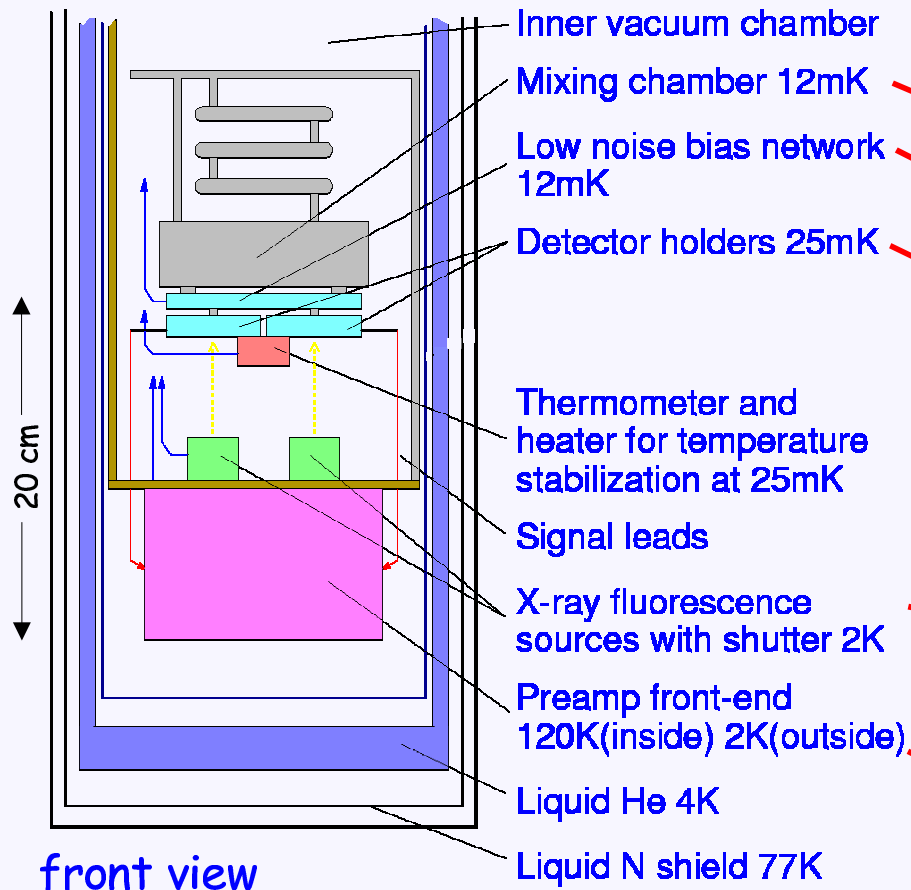
Phase I R&D

- apply and improve silicon micromachined devices (collaboration with **IRST** and **NASA/Wisconsin** groups)
- optimize implant doping and geometry
- improve AgReO_4 crystal quality
- optimize and standardize crystal-thermistor coupling
- improve electronic cryogenic front-end with new JFETs and new design
- develop new gain stabilization techniques
 - ▶ 200 hundred microcalorimeters array
 - ▶ single detector absorber mass increased to $450 \mu\text{g}$
 - ▶ energy resolution ΔE_{FWHM} improved to $10 \div 15 \text{ eV}$
 - ▶ time resolution $\Delta\tau$ improved to $0.2 \div 0.1 \text{ ms}$

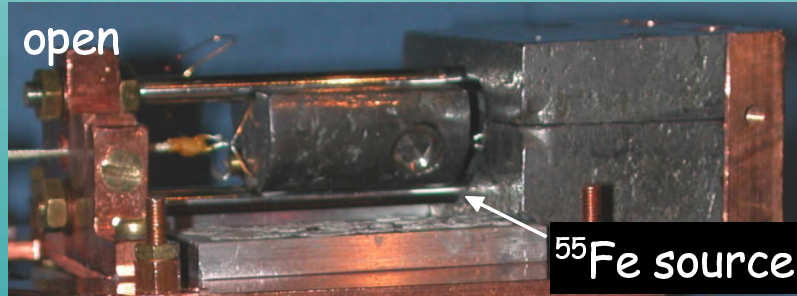
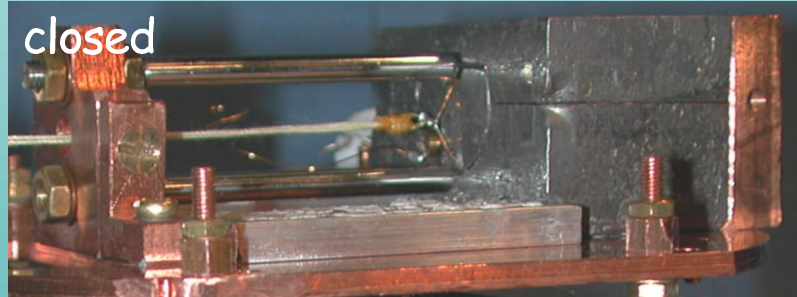
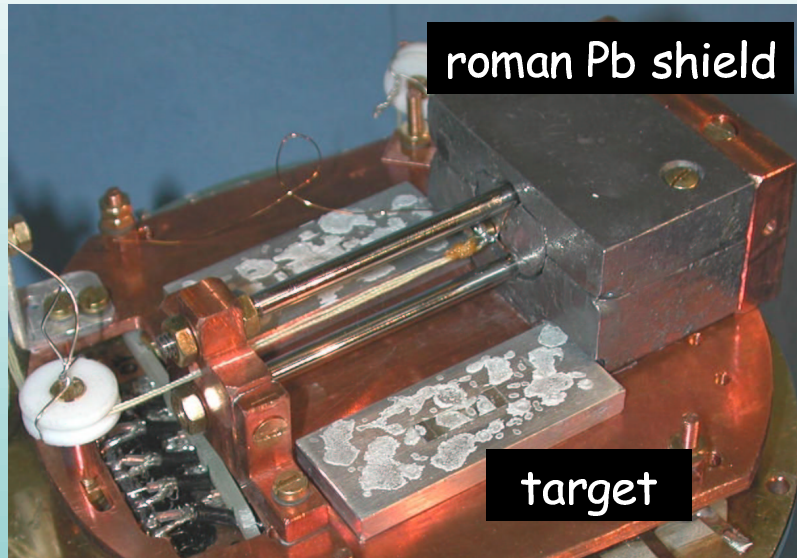


Experimental set-up

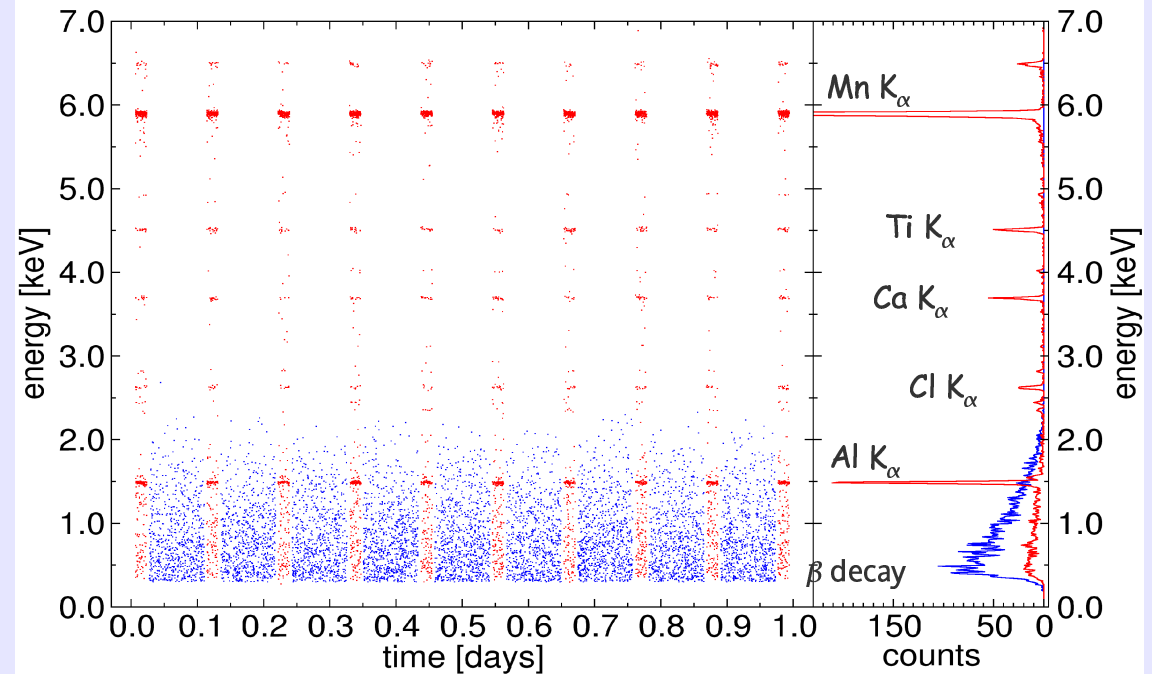
Dilution refrigerator set-up for 10 detector experiment



X-ray fluorescence source for periodic calibration

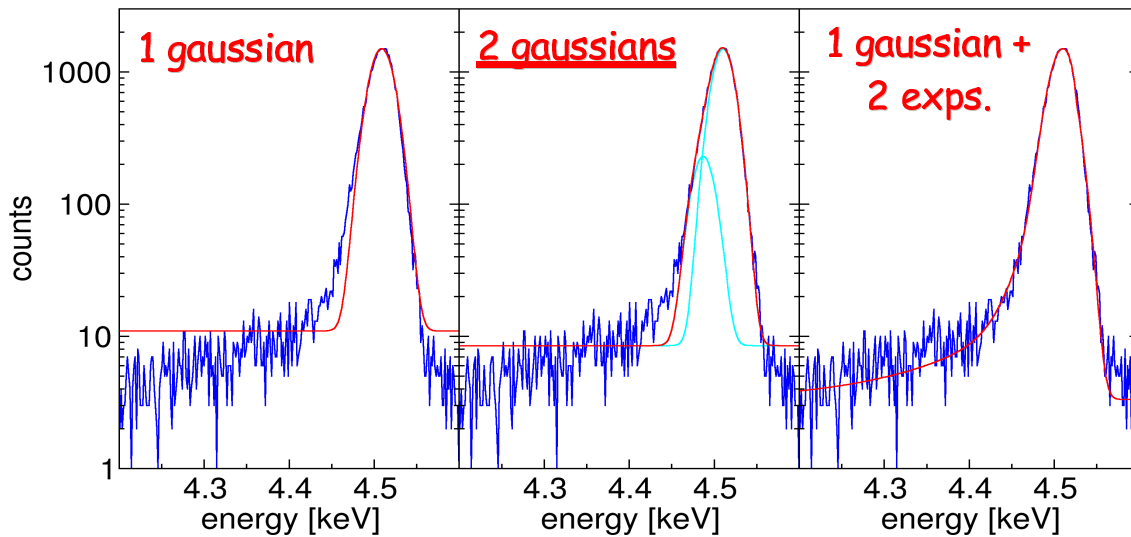


- remotely controlled for continuous stability control
→ 1 cycle: 25 minutes open, 2 hours closed
- primary sources: $2 \times {}^{55}\text{Fe}$ 5 mCi
- roman lead to shield ${}^{55}\text{Fe}$ Inner-Bremsstrahlung
- fluorescence target:
 - Al K_{α} → 1.5 keV
 - Ca K_{α} → 3.7 keV
 - Cl K_{α} → 2.6 keV
 - Ti K_{α} → 4.5 keV



Detector response function

➤ are the response functions for X-rays and for β s from ^{187}Re decay the

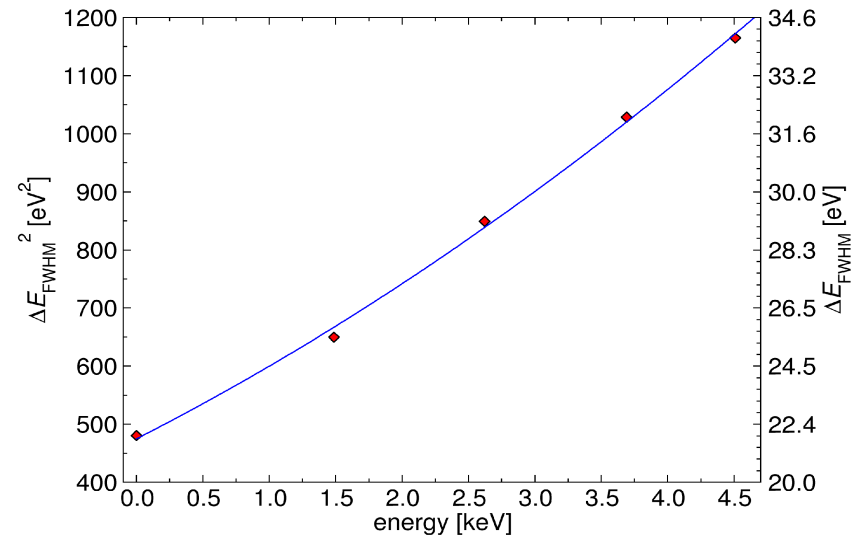


◆ $\lambda(6 \text{ keV}) \approx 3 \mu\text{m}$
 ◆ Absorber dimension $\approx 300 \mu\text{m}$

Ti K_{α} line: $E = 4.511 \text{ keV}$

➤ Energy resolution depends on energy

$\Delta E_{\text{FWHM}} = 28.5 \text{ eV}$
 at $E = 2.466 \text{ keV}$
 (^{187}Re β end-point)



"Internal" calibration of AgReO_4 detectors

- Escape peaks allow internal calibration
 - ▷ $\lambda(6 \text{ keV}) \approx 3 \mu\text{m}$
 - ▷ $\lambda(70 \text{ keV}) \approx 400 \mu\text{m}$
- Escape peaks following incident X-rays are intrinsically broad \Rightarrow seen in previous high statistic measurement with Pb-Re K-K escapes
 - ▷ $\Gamma_{\text{escape KK}} = \Gamma_{\text{PbK}} + \Gamma_{\text{ReK}} \approx 60.2 \text{ eV} + 42.1 \text{ eV} = 102 \text{ eV}$

Internal calibration with ^{44}Ti

- $\rightarrow \gamma$ rays @ 78.4 keV
- \Rightarrow no intrinsic broadening

Re K-edge @ 71.7 keV

γ -X escape peaks have only ReK intrinsic width ($\sim 42 \text{ eV}$)

