

Fig. 2. Two tomograph examples, each one corresponding to a different type of tomograph image. The left tomograph is a CV-tomograph, while the right a CH-tomograph. Both of them are defined by the temporal ordering of lines that pass from the center of the frames. Five indicative frames of the shot from which each tomograph was generated are also shown to the left of the corresponding tomograph. The temporal order of the shown frames and the time-axis of the 2D tomograph images are denoted with arrows.

are combined to generate the final concept detection output. Consequently, assuming that the classification is performed at shot level, the associated complexity is proportional to $S \cdot D \cdot \Lambda$, where S is the total number of classified shots, D is the amount of concepts and Λ is the number of available base detectors.

The two most common late fusion strategies are averaging the confidence scores or calculating a linear combination of them with weights that are globally tuned for all concepts (e.g. [2]). The latter approach, while shown to enhance the concept detection accuracy, suffers from the "curse of dimensionality" that prohibits a brute-force tuning of the weights, especially since the number of base detectors is typically in the order of tens. Moreover, both late fusion approaches do not take into account the fact that not all of the employed detectors can truly contribute to the detection of all possible concepts. Therefore, computational time is unnecessarily consumed, since all base detectors are used in each concept, regardless if they increase the concept detection accuracy or not. Finally, as will be subsequently demonstrated, when the detection accuracy measure depends on the sorting of the results (as is the case with Average Precision (AP) or Extended Inferred Average Precision (XInfAP) [34]), averaging the confidence scores of two base detectors out of which only one achieves good concept detection accuracy may lead to the accuracy of the final detector (fused scores) being worse than the accuracy of the best out of the two original base detectors.

In order to clarify the last point, we model the confidence score distribution of a base detector that is trained to detect a specific concept as a mixture of two gaussian components $N(\mu_p, \sigma_p)$, $N(\mu_n, \sigma_n)$, each corresponding to true positive and true negative samples, respectively. The parameters that determine the classifier accuracy are $\mu_p, \sigma_p, \mu_n, \sigma_n$, as well as the concept's prior probability P_p . Since the Average Precision measure depends on the distance between the two gaussian components, the classifier is translation invariant. Thus, μ_n is arbitrarily selected to be equal to 0. Consequently, the performance of a detector C_3 that averages the scores of two base detectors C_1, C_2 is controlled by 7 parameters (parameters

 $\mu_p, \sigma_p, \sigma_n$ for each of C_1, C_2 , and P_p , since the value of the latter depends only on the frequency of the concept in the dataset). In the subsequent analysis we examined 6 different values of P_p , 1%, 5%, 16.6%, 33.3%, 50% and 66.6% in order to model concepts that are rare as well as concepts that can be found very often in a video stream. For each P_p value we estimate the probability of the following hypothesis being true: " $P(AP(C_3) > max(AP(C_1), AP(C_2)))$ " (this probability we denote as P_0 in the sequel). We estimate the value of P_0 as a function of $AP(C_1)$ and $AP(C_2)$ (i.e. the average precision of the two base classifiers being combined), and we compare it with 0.5, since $P_0 > 0.5$ signifies that the case of detector C_3 performing better than the best detector among C_1, C_2 is more probable than the opposite one.

Plotting P_0 exclusively as a function of $AP(C_1)$ and $AP(C_2)$, for a chosen value of P_p , so as to reveal the relationship between P_0 and the AP scores of the base classifiers, is achieved through the following procedure. First, we randomly selected a large number of possible σ_p, σ_n value pairs (10⁶ different pairs in our simulations). Then, for a given value of μ_p , we estimated the AP score of a base detector C_1 (or C_2) for each possible σ_p, σ_n value pair, and the average of these AP scores we considered as the AP score for the chosen value of μ_p (independently of σ_p, σ_n). Repeating this simulation for different values of μ_p , we found that this AP score is monotonically increasing with μ_p , and we built a lookup table allowing us to estimate the expected μ_p value that corresponds to a chosen AP score for C_1 (or C_2). Subsequently, P_0 can be plotted as a function of $AP(C_1)$ and $AP(C_2)$ as follows: (a) for each pair of base detector AP scores, the corresponding pair of μ_p values are found (using the lookup table), (b) a random pair of σ_p, σ_n values is selected, (c) using these parameter values the AP performance of the detector C_3 is estimated and compared with the maximum value of C_1 and C_2 , and d) this process is repeated, similarly to when estimating the AP of a base detector above, for multiple pairs of σ_p, σ_n values. Following this, the value of P_0 is retrieved as the relative frequency of $AP(C_3)$ being greater than $max(AP(C_1), AP(C_2))$.

The results of the above simulation are demonstrated in Fig. 3. In this figure, starting from the top-left corner, the (i,j) block shows the value of P_0 when C_1 and C_2 base detectors have AP equal to 0.15 + 0.05i and 0.15 + 0.05j, respectively. The grey level of each block represents the P_0 value, with white standing for $P_0 = 1$ and black for $P_0 = 0$. Moreover, a dot is drawn in the center of block (i,j) if $P_0(i,j) > 0.5$. It should be noted that we have chosen this method of visualization instead of reporting the exact P_0 scores, since the process we described above and used for getting these results is an approximate simulation for extracting mostly qualitative conclusions, rather than an exact mathematical model for the accurate quantitative analysis of AP. Fig. 3 demonstrates that the common assumption that the use of more classifiers always increases the accuracy of their combination is not to be taken as a general rule. On the contrary, if the detection performance already reached by a base detector is relatively high, combining additional base detectors with it or not needs to be thoroughly examined in order to make sure that the accuracy of the combination will not deteriorate, compared to using the single well-performing detector alone. More importantly, if an extensive set of base detectors is available for multiple concepts, a procedure for selecting for each concept the optimal subset of base detectors to be combined is needed.

Moreover, Fig. 3 manifests that P_0 exhibits a different behavior for concepts having different prior probabilities. Specifically, a significant difference between rare and frequent concepts is that in the first case accurate base detectors "dominate" over all other base detectors, since their combination enhances the performance even when the other base detector exhibits a low AP (e.g. for $P_p=1\%$ or $P_p=5\%$). On the contrary, when P_p increases, then the poor base detectors "dominate" the combination, since the performance remains low even when the other base detector of the combination exhibits a high AP. This point explains the disagreement in conclusions between participants in the TRECVID competition (that includes many rare concepts), who have found employing multiple base detectors and late fusion advantageous, and other works that, following the analysis of concept detection results for mostly frequent concepts, rejected such a late fusion approach (e.g. [24], [35]).

B. Base detector selection for concept detection

Motivated by the above analysis, we have developed a base detector selection procedure that selects for each concept the optimal subset of the available base detectors. More specifically, the introduced scheme builds upon a genetic algorithm, by post-processing the outcome of two different variations of the latter. We should stress that this algorithm is executed off-line, during training. The result of this technique is the selection for each concept of an optimal subset of base detectors and the exclusion of all other base detectors from the corresponding detection scheme. Thus, at run-time, the computational complexity of concept detection is reduced from $O(S \cdot D \cdot \Lambda)$ (Section IV-A) to $O(S \sum_{i=1}^D M_i)$, where M_i is the number of base detectors employed for the detection of the i-th concept.

The genetic algorithm that plays a central role in this approach is summarized in Alg. 1. In the following, we use the operator # to represent set cardinality, and the \circ operator to represent the vector Hadamard product (i.e. the result of element-wise multiplication). Moreover, we denote L the ordered set of base detectors, L_i a subset of this set, p_i the AP achieved by using the average of the base detectors score that belong to L_i and v_i the participation vector of subset L_i . As participation vector of a subset L_i we refer to a binary vector of length #L, whose j-th element is equal to 1 if and only if the j-th element of L belongs to L_i .

Algorithm 1 Genetic algorithm for selecting a subset of best-performing (or worst-performing) base detectors.

Notation: c is the current concept, m the mutation rate of the genetic algorithm, N is the initial population size, R the (fixed) number of generations and k is the number of parent chromosomes that breed the next generation population.

- 1: Initially, from set L, N random subsets (chromosomes) $L_1, L_2, ..., L_N$ are used to form the initial population. Their corresponding participation vectors $v_1, v_2, ..., v_N$, as well as the corresponding performance estimations $p_1, p_2, ..., p_N$ are computed. The current generation index r is set to 1.
- 2: The k chromosomes that achieved the best (or worst) performance (k < N) "survive", while all the other chromosomes are discarded.
- 3: Uniform crossover is used to combine the k parent chromosomes of the current generation in k(k-1)/2 pairs to breed two new chromosomes each, thus leading to a new population of k(k-1) members. More specifically, from two parent chromosomes L_i and L_j the children chromosomes will have participation vectors $v_i \circ v_j + Y \circ (1 v_i \circ v_j)$ and $v_i \circ v_j + (1 Y) \circ (1 v_i \circ v_j)$, where Y is a random binary vector of dimension #L, $\sum Y = \#L/2$, and "1" denotes a vector of ones. Thus, each child chromosome inherits some of its genes from both its parents.
- 4: Once the new population is constructed mutation is employed to randomly modify a gene subset. More specifically, out of the k(k-1)(#L) genes of the population, mk(k-1)(#L) of them randomly mutate (i.e. the corresponding participation vector elements change value from 0 to 1 or from 1 to 0).
- 5: The chromosomes that match the k(k-1) participation vectors formed after the end of step 4 are retrieved and the corresponding performance is estimated.
- 6: If r=R then the chromosome L_0 that achieved the maximum performance is returned as the optimal configuration. Moreover, the participation vectors $v_1, v_2, ..., v_T$ of the chromosomes that achieved the top-T performance values are retrieved. Otherwise, r=r+1 and the algorithm continues from step 2.

The aforementioned genetic algorithm is executed independently twice under the proposed approach. In the first execution the goal is to identify the configurations (i.e. the base detector subsets) that achieve the best performance, while in the second one the configurations that exhibit the worst performance. These are used for robustly selecting the best base

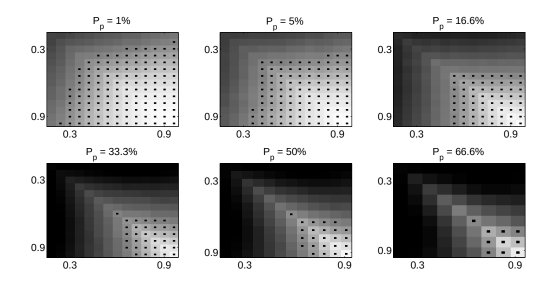


Fig. 3. The probability of a classifier that averages two base detectors to exhibit higher AP performance than the best-performing one of these base detectors, as a function of the AP of the two base detectors. The two axis correspond to the AP performance of each base detector. More specifically, starting from the top-left corner, the (i,j) block corresponds to a pair of detectors, having AP equal to 0.15 + 0.05i and 0.15 + 0.05j, respectively. The grey level of each block represents the value of probability P_0 , with white standing for 1 and black for 0. Moreover, a black dot in the center of a block (i,j) denotes that $P_0(i,j) > 0.5$.

detectors for the given concept: a base detector is ultimately selected if it is frequently included in the best-performing configurations and at the same time is not frequently included in the worst-performing ones. For realizing this selection, we introduce a "base detector quality" measure Q_c , in relation with concept c,

$$Q_c = (P_T - N_T)/T \tag{4}$$

where T is the number of configurations that are retrieved in each genetic algorithm execution, and P_T, N_T is the number of times that the base detector was included in the configurations that achieved the top-T and the bottom-T performance, respectively. The base detectors for concept c are ranked using Eq. (4), and the M highest-ranked ones are selected. The above described process is summarized in Alg. 2.

It should be noted that fixing the number of base detectors M_i that are selected for each concept to $M_i=M$, as we do in step 4 of Alg. 2, is a choice we make for simplifying the experimental evaluation and comparison of the proposed approach. In practice, one could also search for the optimal number of base detectors separately for each concept, which may further improve the results.

V. EXPERIMENTAL EVALUATION

A. Datasets and experimental setup

Our experimental setup is based on the 2011 and 2012 TRECVID SIN Tasks [36], [37]. As already mentioned, the total number of concepts that were defined in these tasks is 346. However, the corresponding evaluations were carried out in a subset of 50 and 46 concepts, respectively, for which ground-truth annotation of the test sets is available. The latter concepts, which are the ones used in this work, are shown in Table I. In this table we additionally mark with a "*" the concepts that are either directly related with motion (e.g.

Algorithm 2 The proposed base detector selection algorithm.

- 1: For each concept the genetic algorithm (Alg. 1) is executed. In each generation of the algorithm the configurations that achieved the best performance "survive". The output is the top-T configurations that achieved the best performance for each concept and the mean base detector number M (i.e. the average of the number of base detectors in the top-1 configurations across all concepts, rounded to the nearest integer).
- 2: For each concept the genetic algorithm (Alg. 1) is executed again. This time, in each generation of the algorithm the configurations that achieved the worst performance "survive". The output is the bottom-T configurations that achieved the worst performance for each concept.
- 3: For each concept and each base detector, the number of times that this base detector was included in the top-T and the bottom-T configurations (i.e. P_T and N_T , respectively) is estimated, and the base detector quality measure Q_c is calculated according to Eq. 4.
- 4: For each concept the M base detectors with the highest Q_c comprise the base detector subset (i.e. configuration) that will be employed in the concept classifier, while all other base detectors are discarded.

"throwing", "walking-running", "skating") or correspond to objects that are very likely to be filmed while they are in motion (e.g. "skier", "car", "boat-ship").

For training, validation and testing our concept detectors we have used the TRECVID SIN Task datasets as follows: For the experiments on the 2011 dataset our training set consisted of 11644 videos (lasting approximately 400 hours and including 250000 shots), the validation set of 4216 videos (approx. 100 hours; 70000 shots) and the test set of 4000 videos (approx.

TABLE I
The 50 and 46 concepts evaluated in the 2011 and 2012 TRECVID SIN dataset, respectively. Motion-related concepts are marked with a "*".

Dataset	Concepts						
TRECVID 2011	Adult, Anchor-person, Beach, Car*, Charts, Cheering*, Dancing*, Demonstration*, Doorway, Explosion/Fire,						
	Face, Female Person, Female Face Close-up, Flowers, Hand, Indoor, Male Person, Mountain, News Studio,						
	Nighttime, Old People, Overlaid Text, People Marching*, Reporters, Running*, Scene Text, Singing, Sitting						
	Down*, Sky, Sports*, Streets, Two People, Walking*, Walking/Running*, Door Opening*, Event*, Female Human						
	Face, Flags, Head & Shoulder, Male Human Face, News, Quadruped, Skating*, Speaking, Speaking To Camera,						
	Studio With Anchor-person, Table, Text, Traffic*, Urban Scenes						
TRECVID 2012	Airplane*, Airplane Flying*, Basketball*, Bicycling*, Boat-Ship*, Boy, Bridges, Chair, Computers, Female						
	Person, Girl, Government leader, Greeting*, Highway*, Instrumental Musician, Kitchen, Landscape, Male Person,						
	Meeting, Motorcycle*, Nighttime, Office, Press Conference, Roadway Junction, Scene Text, Singing, Sitting						
	Down*, Stadium, Teenagers, Throwing*, Walking-Running*, Apartments, Baby, Civilian Person, Clearing, Fields,						
	Forest, George Bush, Glasses, Hill, Lakes, Man Wearing A Suit, Military Airplane*, Ocean, Skier*, Soldiers*						

100 hours; 65000 shots). For the experiments on the 2012 dataset, the training set consisted of 19860 videos (approx. 600 hours; 400000 shots), the validation set of 4163 videos (approx. 100 hours; 73000 shots), while the test set of 4100 videos (approx. 100 hours; 72000 shots).

For evaluating the trained concept detectors we followed the methodology used in TRECVID. That is, for each concept separately, the top 2000 shots sorted by detection score in descending order are returned and are evaluated against partial manually-generated ground-truth annotations. The evaluation measure is the Extended Mean Inferred Average Precision (XInfAP) [34], which has been proposed for the purpose of approximating Average Precision (AP) when the dataset is not fully annotated and therefore AP cannot be directly calculated.

The proposed concept detection approach is implemented, for the purpose of experimental evaluation, according to the pipeline of Fig. 1, employing state-of-the-art features and parameters for the pipeline components that are not explicitly discussed in the preceding sections. Specifically, each video shot is represented by either one or more keyframes or also a pair of video tomographs (CV-tomograph, CH-tomograph). Subsequently, an interest point detector is employed to select the image points at which descriptors will be extracted. We used two such detectors; the first selects interest points through dense sampling, i.e. in fixed distances in a 2D image grid, while the second one is a Harris-Laplace corner detector [8]. At each of the resulting interest point locations, lowlevel visual descriptors are extracted (SIFT, RGB-SIFT and Opponent-SIFT), following the conclusions drawn in [5] for video concept detection tasks. Subsequently, the low-level descriptors are assigned to visual words, using two vocabularies that were created off-line through k-means clustering and hard- or soft-assignment, respectively, according to [38]. In all cases (i.e. regardless of which one of the above video sampling strategies, descriptors etc. is used) a pyramidal 3×1 decomposition scheme employing 3 equally-sized horizontal bands of the image, as proposed in [39], was used on every keyframe or tomograph, generating 3 different BoWs corresponding to the three image bands and a fourth BoW for the entire image. The number of words for each BoW was set to 1000 and the four BoWs coming from the adopted pyramidal decomposition were concatenated to a 4000-element BoW vector. One such vector is calculated separately for

each combination of representation (i.e. each keyframe or tomograph), interest point detector, descriptor and assignment method and is used as input to the corresponding SVM base classifiers for a given concept (base detectors). As a result, 36 base detectors are built for each concept separately, based on local image features. An additional base detector that uses a global visual descriptor (HSV histogram) is also employed. The 37 base detectors are outlined in Table II.

For the base detectors, linear SVM classifiers are employed instead of the kernel SVMs that are typically used in such tasks. By this choice, the required computational time for a single SVM classifier (corresponding to a single concept) fell from 6 seconds per image (that was required in our earlier experiments with kernel SVMs) to 0.03 seconds. This is an implementation choice that is in line with our overall goal of developing a computationally efficient solution to concept detection. All base detectors were trained off-line, using the corresponding training sets. The output of each of the trained base detectors is an intermediate confidence score. The overall confidence score for each concept is estimated as the harmonic mean value of the intermediate confidence scores of all (i.e. up to 37) base detectors for this concept. Using the harmonic mean was shown in practice to produce slightly better results than the arithmetic mean, which was used for simplicity in our simulations reported in Section IV-A.

B. Tomographs versus keyframes

In this first experiment we investigate the impact of using the video tomographs introduced in Section III in replacement of traditional keyframes. More specifically, we compare the combination of the 13 keyframe-based detectors of Table II against the combination of 12 tomograph-based detectors of the same Table that use dense sampling for interest point detection (which, as we show later on, are the best among our 24 tomograph-based detectors). The results show that out of the 96 different concepts in the two datasets that we use, tomographs outperform keyframes for only 8 concepts (all of them motion-related: basketball, throwing, skier, demonstration, people marching, running, walking/running, skating). This result is consistent with our intuition that tomographs convey useful motion information that is, however, complementary rather than alternate to the non-motion information

TABLE II
COMPLETE SET OF BASE DETECTORS USED IN OUR EXPERIMENTS.

Video Sampling	Possible representations and resulting base detectors						
Keyframes	12 local-image-feature-based Base Detectors: 3 descriptors (SIFT, Opponent-SIFT, RGB-SIFT) × 2 point						
	detectors (Dense, Harris-Laplace) × 2 BoW strategies (soft-, hard-assignment)						
	1 global-image-feature-based base detector (HSV color histograms)						
Tomographs	24 tomograph-based base detectors: 2 types of video tomographs (CH-tomograph, CV-tomograph) × 3						
	descriptors (SIFT, Opponent-SIFT, RGB-SIFT) ×2 point detectors (Dense, Harris-Laplace) × 2 BoW						
	strategies (soft-, hard-assignment)						

conveyed by keyframes. Based on this result and in accordance with the emphasis that we put in all previous sections on how to combine tomographs and keyframes, we will proceed in the sequel with experimentally evaluating such combinations.

C. Combining tomographs and keyframes

In this series of experiments we investigate the impact of combining the video tomographs introduced in Section III with traditional keyframes. The following approaches are compared: i) A13 is the baseline, using 1 keyframe per shot and combining the 13 keyframe-based base detectors of Table II, ii) A25 combines the 13 base detectors of A13 with 12 additional base detectors that employ tomographs as video samples and use dense sampling for interest point detection, iii) A37 combines all 37 base detectors of Table II, iv) A39 employs the same base detectors as A13, but, as proposed in [3], in this case 3 keyframes per shot are used instead of one and therefore the 13 base detectors of A13 have to be evaluated 3 times each for a single shot (thus, 39 base detector scores are produced per shot). Our choice to compare tomographs with the multiple keyframe approach of [3] is based on the fact that any other existing solution (e.g. traditional motion descriptors) would feature significantly higher computational cost, making the application of it on the extensive TRECVID datasets difficult and in any case depriving such a comparison of any practical value.

The experimental results are summarized in the top block of Table III (MXInfAP, i.e. mean XInfAP across all concepts). Comparing A25 with A13, the MXInfAP increases in the 2011 dataset from 0.2187 to 0.2659 and in the 2012 dataset from 0.1394 to 0.1557. These improvements represent a 21.6% and 11.7% accuracy boost, respectively, and manifest that although the tomographs are not potential replacements of traditional keyframes, they provide additional information that the latter do not capture. Moreover, as can be seen in Table III, it is the dynamic (motion-based) concepts that benefit the most from the introduction of tomographs: if only the motion-related concepts are taken into account, then the accuracy boost caused by introducing tomographs is 38.9% and 47.1% for the two testsets, respectively.

On the other hand, the reported results show that the detection of interest points by means of Harris-Laplace in tomograph images generates noisy representations that lower the overall performance. This is manifested by the significantly lower performance that A37 exhibits compared to A25 (MX-InfAP of 0.2113 versus 0.2659 and 0.0837 versus 0.1557 in the 2011 and 2012 datasets, respectively). This is intuitively

supported by the fact that tomographs are by default very noisy images, thus selecting image corners as interest points in such images does not necessarily help. However, as we show in the next subsection, if our concept detection problem focuses on motion-related concepts and base detector selection can be performed for each concept, then base detectors of this type can also contribute to more accurate results.

We also compare A25 with A39, which performs a denser video sampling in the temporal direction [3], thus using a larger number of keyframes for each shot. While A39 employs 13 distinct base detectors (the same used in A13), the use of 3 keyframes per shot results in each base detector being evaluated 3 times for each shot and therefore needs computational time similar to that of A37 (we should note that "computational time" here and throughout the rest of the Experimental Evaluation section refers to the time needed for obtaining a classification result for a given non-classified shot). Despite the additional keyframes, the performance of A39 is significantly lower compared to that of A25, while the latter is also less computationally demanding (Table III). This shows that tomographs should be preferred over introducing additional keyframes in a keyframe-based concept detection scheme.

Finally, in Figs. 4 and 5 we show results per concept (XInfAP) for the A13, A25 and A37 experiments. Although many of the considered concepts are not intuitively expected to be strongly correlated with any type of motion (e.g. "landscape", "fields", "computers"), we can see that when combining keyframe-based and certain tomograph-based base detectors (A25), XInfAP increased for more than 80% of the concepts.

D. Selecting base detectors for building a concept classifier

In this series of experiments we investigate the impact of using a base detector selection strategy for choosing a different subset of base detectors for each concept that we want to detect. For this we apply the technique introduced in Section IV and evaluate it against the results of the previous set of experiments (where no such selection strategy was used) as well as against variations of the technique of Section IV and related literature works [13], [40], [25]. More specifically, starting from the 25 base detectors used in A25, the following approaches are compared: i) G25a, a variation of the proposed technique in which only the first step of Alg. 1 is executed and the chromosome from the initial population whose performance is maximum at the end of that step is returned as the chosen configuration (i.e. set of base detectors), ii) G25b,

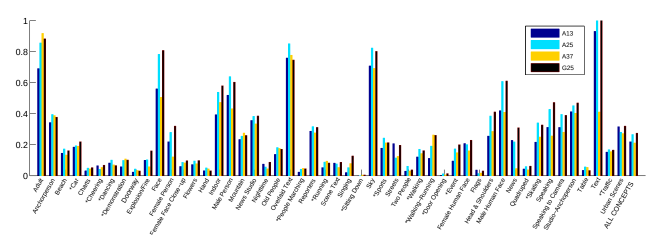


Fig. 4. XIinfAP for the concepts of the 2011 dataset. The compared techniques are A13, A25, A37 and G25.

TABLE III EXPERIMENTAL RESULTS AND COMPARISONS. AVERAGE BOOST IS THE RELATIVE PERFORMANCE INCREASE COMPARED TO A13, while # Base Detectors is the average number of base detectors per concept and shot that need to be evaluated at run time.

•	TRECVID 2011 dataset				TRECVID 2012 dataset					
	All co	ncepts		-related		All concepts		Motion-related		
			concep	ts only				concepts only		
Method	MX-	Aver-	MX-	Aver-	# Base	MX-	Aver-	MX-	Aver-	# Base
	InfAP	age	InfAP	age	Detec-	InfAP	age	InfAP	age	Detec-
		Boost		Boost	tors		Boost		Boost	tors
Tomographs at	nd keyframe	es vs kefra	mes only (experimen	ts of Section	on V-C)				
A13	0.2187	0%	0.0959	0%	13	0.1394	0%	0.0797	0%	13
(baseline)										
A25	0.2659	21.6%	0.1332	38.9%	25	0.1557	11.7%	0.1172	47.1%	25
A37	0.2113	-3.4%	0.1247	30%	37	0.0837	-39.7%	0.0845	6%	37
A39	0.2417	10.5%	0.1126	17.4%	39	0.1471	5.5%	0.0911	14.3%	39
Base detectors	selection a	mong 25 b	ase detecto	ors (experi	ments of S	ection V-D))			
G25a	0.2536	16%	0.1291	34.6%	10.32	0.1571	12.7%	0.1177	47.7%	10.11
G25b	0.2701	23.5%	0.1339	39.6%	10.02	0.1651	18.4%	0.1238	55.3%	8.89
G25	0.2744	25.5%	0.1346	40.4%	10	0.1783	27.9%	0.1299	63%	9
R25 [13]	0.2247	2.7%	0.1197	24.8%	25	0.1422	2%	0.1023	28.4%	25
ABC25 [40]	0.2617	19.7%	0.1282	33.7%	10.52	0.1697	21.7%	0.1282	60.9%	8.87
GA25 [25]	0.2585	18.2%	0.1301	35.7%	25	0.1667	19.6%	0.1278	60.4%	25
Base detectors	selection a	mong 37 b	ase detecto	ors (experi	ments of S	ection V-D)		L	
G37a	0.2411	10.2%	0.1283	33.8%	13.02	0.1506	8%	0.1217	52.7%	13.69
G37b	0.2633	20.4%	0.1307	36.3%	12.74	0.1659	19%	0.1329	66.8%	11.78
G37	0.2743	25.4%	0.138	43.9%	13	0.1760	26.3%	0.1361	70.8%	12
R37 [13]	0.2317	5.9%	0.1202	25.3%	37	0.1387	-0.5%	0.0983	23.3%	37
ABC37 [40]	0.2635	20.5%	0.1274	32.9%	12.96	0.1639	17.6%	0.1296	62.6%	12.16
GA37 [25]	0.2648	21.1%	0.134	39.7%	37	0.1681	20.6%	0.1341	68.3%	37

another variation of the proposed technique in which only the first step of Alg. 2 (i.e. the complete Alg. 1) is executed and the chromosome whose performance is maximum at the end of that step is returned as the chosen configuration, iii) G25, which corresponds to the complete approach proposed in Section IV-B, iv) ABC25, which uses the Artificial Bee Colony (ABC) [40] algorithm for base detector selection, v) R25, in which the 25 available confidence scores are linearly combined using weights determined through regression, as proposed in [13], and vi) GA25, in which the 25 available confidence scores are linearly combined using weights determined through gradient ascent, as proposed in [25]. All approaches are executed independently for each concept of both the 2011 and 2012 TRECVID datasets. Subsequently, the above 6 experiments are repeated, this time starting from the 37 base detectors used in A37 (thus, 37 base detectors are

available for selecting a subset of them, as opposed to 25 in the previous 6 experiments). These experiments are denoted G37a, G37a, G37, ABC37, R37, GA37, respectively. For running the above experiments, the Alg. 1 parameters were set to N=5000, R=60, k=40, m=0.15 and T=500, while the employed Artificial Bee Colony (ABC) parameters were the ones proposed in [40].

The results of the experiments (MXInfAP) are reported in the second and third block of Table III, together with the corresponding average performance boost compared to A13 and the mean number of base detectors that were used in each experiment. The latter number is assumed to be approximately proportional to the computational cost of the overall concept detection pipeline at run time.

Firstly, the results show that the G25 and G37 approaches compare favorably to the A13, A25 and A37 ones examined

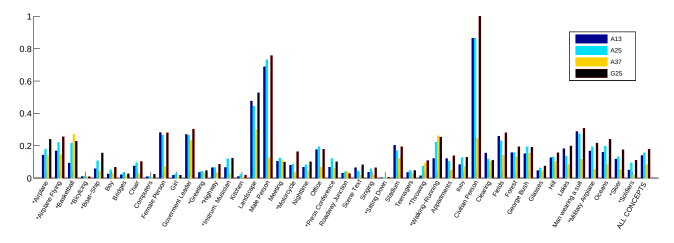


Fig. 5. XIinfAP for the concepts of the 2012 dataset. The compared techniques are A13, A25, A37 and G25.

in Section V-C, thus demonstrating the impact of the proposed base detector selection strategy to both MXInfAP and computational cost. Secondly, we can see that the results of G25and G37 are very similar (the MXInfAP difference between G25 and G37 is only 0.0001 and 0.0023 in the 2011 and 2012 dataset). Following the discussion in Section V-C on how Harris-Laplace interest point detection in tomographs leads to noisy base detectors, this similarity of results shows that the proposed base detector selection strategy can effectively identify and discard noisy detectors. Further to this, we can see in Table III that when considering motion-related concepts only, G37 outperforms G25; this demonstrates the merit of even noisy base detectors under certain conditions. G25 is further compared with A13, A25 and A37 for each concept separately, and the results are shown in Figs. 4 and 5, where it can be seen that G25 improves on the results of A13, A25and A37 for the vast majority of the concepts.

Comparison of G25 and G37 with simpler variants of them, namely G25a, G25b, G37a, G37b (Table III), shows the significance of proposed Algs. 1 and 2 in making a stable selection of good base detectors for each concept. Particularly the importance of taking into account not just the top-T but also the bottom-T configurations when selecting base detectors (second step of Alg. 2) is made clear by this comparison.

With respect to other SoA methods (Table III), the proposed G25 (or G37) approach compares favorably to ABC25 [40] (or ABC37), a metaheuristic search algorithm for optimization. This can be explained by the fact that the ABC technique does not take into account that in the concept detection task the solution space is noisy, thus the optimization in a validation set often does not lead to good detection in the testset. Similar conclusions can be drawn by comparing G25 (or G37) with GA25 [25] (or GA37) and R25 [13] (or R37), the most recent base detector combination approaches that can be found in the relevant literature. It seems that the performance reached through gradient ascent (i.e. GA25) is similar to the performance reached through a simple genetic algorithm (G25b) or ABC (ABC25). However, linear combination approaches by definition use the confidence scores of

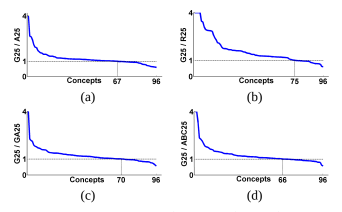


Fig. 6. Comparison of G25 against A25, R25, GA25 and ABC25 at the individual concept level, using the ratio of XInfAP achieved by G25 and (a) A25, (b) R25, (c) GA25 and (d) ABC25. In each sub-figure, the XInfAP ratio is estimated for each of the 96 concepts and then used for sorting the concepts in descending order. Ratios higher than 4 are truncated to 4 for visualization purposes. A dotted horizontal line signifies a ratio equal to 1, while the number of concepts with ratio higher than 1 is highlighted on the x-axis.

all base detectors. Consequently, both GA25 (GA37) and R25 (R37) are much more computationally demanding at runtime than the corresponding algorithms that use a base detector selection scheme (e.g. G25). Finally, the poor results of R25 (R37) can be explained by the fact that for most concepts the number of positive samples in the dataset is much lower than the number of negative samples. This bias undermines this technique, which minimizes the total sum of the error without discriminating between positive and negative samples.

Further comparison of G25 against A25, R25, GA25 and ABC25 at the individual concept level (Fig. 6) shows that the former outperforms the latter for 67, 75, 70 and 66 of the 96 concepts, respectively (and is outperformed by only a small margin for most of the few remaining concepts). These results manifest that, although the proposed optimization approach does not theoretically guarantee optimal base detector selection (as does neither one of the related literature approaches), in practice it consistently performs well in these 96 optimization problems.

E. Assessing the impact of algorithm parameters and design choices

A critical parameter of the approach of Section IV is the number of base detectors that are selected for each classifier. We have plotted the MXInfAP of G25 and G37 as a function of this number (Fig. 7). The plot demonstrates that the achieved MXInfAP does not significantly depend on the exact base detector number, since in all cases there is a rather wide area around its optimal value where it remains almost constant. More specifically, in the G25 case the optimal base detector number is 13 in the 2011 dataset and 8 in the 2012 one. The proposed strategy, which is to automatically select the number of base detectors by the genetic algorithm outcome, leads to the selection of 10 and 9 base detectors, respectively. The MXInfAP loss as a result of this is 0.006 in the 2011 dataset (since 0.2804 is the theoretical maximum as seen in Fig. 7, and 0.2744 is the one actually achieved) and 0.0005 in the 2012 dataset (0.1788 and 0.1783 respectively). In the G37 case, for both 2011 and 2012 datasets the optimal base detector number is equal to 13, while the proposed strategy leads to the selection of 13 base detectors in the 2011 dataset and 12 base detectors in the 2012 dataset. In the second case, the corresponding MXInfAP loss is 0.0012 (from 0.1772 to 0.176).

For completeness, it should be noted that if a variable number of base detectors is used for each concept, then the overall MXInfAP may further increase. In experiments where we varied the value of M_i between 6 and 25 separately for each concept and selected the optimal configurations, the maximum achieved MXInfAP was 0.2938 and 0.1858 for the 2011 and 2012 dataset, respectively (representing an up to 7% boost over the G25 results reported in Table III).

In order to validate our design choice of defining the base detector quality Q_c according to Eq. 4, for selecting with the help of the genetic algorithm the optimal set of base detectors, we further examine the impact of using, in place of Q_c , one of three other reasonable base detector quality measures Q_c^1 , Q_c^2 , Q_c^3 , defined as follows:

$$Q_c^1 = AP_T^P - AP_T^N (5)$$

$$Q_c^2 = P_T/T \tag{6}$$

$$Q_c^3 = AP_T^P \tag{7}$$

where P_T (respectively, N_T) is the number of times that the base detector was included in the configurations that achieved the top-T (bottom-T) performance, and AP_T^P (respectively, AP_T^N) is the Average Precision value that can be calculated by treating the list of top-T (bottom-T) configurations as a ranked list of retrieval results, where configurations that include the base detector in question are taken as positive samples. The results, demonstrated in Table IV, manifest the validity of a genetic algorithm that looks at not only the top-T but also the bottom-T configurations, since Q_c and Q_c^1 outperform the corresponding quality measures that

TABLE IV
PERFORMANCE COMPARISON WHEN EACH OF THE FOUR DIFFERENT
QUALITY MEASURES DEFINED IN EQS. 4 - 7 ARE USED.

Method	Dataset	Q_c (4)	Q_c^1 (5)	Q_c^2 (6)	Q_c^3 (7)
G25	2011	0.2744	0.2731	0.2736	0.2728
G25	2012	0.1783	0.1762	0.1691	0.1692
G37	2011	0.2743	0.2714	0.2718	0.2705
G37	2012	0.1760	0.1762	0.1706	0.1680

employ only the top-T configurations, Q_c^2 , Q_c^3 . Additionally, averaging the results (Q_c, Q_c^2) seems to exhibit more stable performance than taking into account the exact rank by calculating some sort of Average Precision (Q_c^1, Q_c^3) , although in general all observed differences are rather small, suggesting that the proposed genetic algorithm is relatively insensitive to the exact way in which Q_c is defined.

F. Statistics on the usefulness of different base detectors

Figure 8 shows the number of times that each base detector is used in all 96 concept classifiers if the G37 approach is followed. It can be seen that the keyframe-based base detectors are selected more often than the tomograph-based ones. Specifically, on average only 33.8% of the 37 available base detectors are used for a given concept, and 10 out of 13 keyframe-based detectors are consistently selected for more than 50% of the examined concepts. There are also 3 tomograph-based detectors that are selected for the majority of the concepts, despite the fact that motion-related concepts are a minority in our datasets (28 motion-related concepts out of 96 concepts in total). Finally, the tomograph-based base detectors that employ interest point detection via Harris-Laplace rather than extracting local descriptors on a dense grid are the least contributing ones, being ranked in the last 12 places in Fig. 8.

G. Discussion on computational complexity

For comparing the methods' time efficiency throughout the paper, we made the assumption (in Section V-D) that the computational cost of the overall concept detection pipeline at run time is approximately proportional to the mean number of employed base detectors. However, under the G25 (and G37) approaches, since a different subset of base detectors is used by each concept classifier, features will in any case need to be extracted for all 25 (or 37) possible base detectors when working with multiple concepts. Thus, the above assumption will only hold if the most computationally expensive step of the pipeline is evaluating the trained SVMs, rather than performing feature extraction and BoW creation. Our experiments show that this is indeed the case under the typical application scenario that involves the detection of a plurality of different concepts, e.g. one thousand or more as in [41], [42]. Specifically, the time needed for evaluating the trained SVMs when detecting one thousand concepts under the A13 and G25approaches (with 9 base detectors being used for each concept in G25) accounts for 97% and 93.9% of the overall processing time, respectively, despite using linear SVMs, as a result of the high number of such SVMs that need to be evaluated. In this experiment, G25 is found to be overall 28.6% faster than A13,

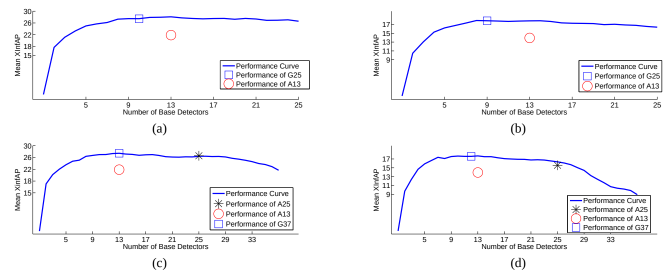


Fig. 7. Performance as a function of the number of selected base detectors. (a) G25 in the 2011 dataset (b) G25 in the 2012 dataset. (c) G37 in the 2011 dataset (d) G37 in the 2012 dataset. For better readability, the numbering of the vertical axes shows only the fractional part of the MXInfAP value, i.e. the integer part "0." of MXInfAP is omitted.

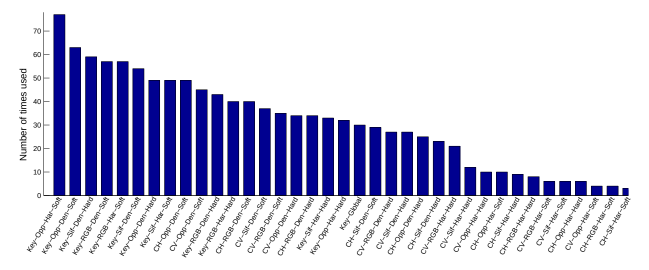


Fig. 8. Barplot of the number of times each base detector is used in all 96 concept classifiers of the two employed datasets, when selection of base detectors is performed according to G37. The naming convention for the base detectors (horizontal axis) is <Video sampling>-<Descriptor>-<Interest point detector>-<BoW assignment strategy>. "Key-Global" refers to the HSV histogram-based base detector. See Table II for details on the base detectors.

as a result of using 30.8% fewer base detectors per concept, despite feature extraction and BoW creation in G25 requiring somewhat more time than in A13.

These run time efficiency gains of base detector selection come at a cost during the training phase, which however is performed off-line and only once for each concept, thus not affecting the scalability of the developed detectors, i.e. their applicability to extremely large volumes of non-labeled video data. Moreover, using the proposed genetic algorithm the training cost is limited because this algorithm visits only a very small portion of all possible configurations. Indicatively, in the most extensive base detector setup (G37) the number of possible configurations is 13.7 billion (i.e. 2^{37}), while the configurations that were examined by the introduced algorithm during our G37 experiment (in all stages) were approximately 100 thousand. Examining a single configuration using our

MATLAB implementation of the base detector selection algorithm on an INTEL(R) Core i7-3770K 3.5GHz PC took 0.7 sec., leading to a total training time of less than 20 hours for each concept; this is certainly not prohibitive for an off-line training process that needs to be performed only once.

VI. CONCLUSION

In this work we dealt with large-scale video concept detection. We showed that video tomographs can contribute to increased concept detection accuracy, both for motion-related and non-motion-related concepts, while introducing no greater computational cost than that of processing a single keyframe. Additionally, we showed that the "one size fits all" approach for the detection of multiple concepts in video streams leads to the unnecessary evaluation of base detectors that do not always contribute to the detection of a specific concept, thus

compromising accuracy while increasing the computational cost. We addressed this issue with a new approach that builds upon a genetic algorithm to rank base detectors and select only a (different for each concept) subset of them. The reported results demonstrate that by combining our base detector selection technique and video tomographs, concept detection effectiveness can be boosted by 25-28% (as measured by MXInfAP), while at the same time the run-time computational cost of concept detection can be decreased as a result of using up to 23-30% fewer base detectors per concept, compared to indiscriminately using the complete set of keyframe-based base detectors.

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